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EVALUATION OF THE VESICATING PROPERTIES
OF NEUTRALIZED CHEMICAL AGENT IDENTIFICATION SET
(CAIS) COMPONENTS

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13. ABSTRACT (Maximum 200 words)

Vesication and skin irritation studies were conducted in hairless guinea pigs to ascertain the vesicant and skin irritation potential of chemically-neutralized Chemical Agent Identification Sets (CAIS). CAIS are training items that contain agent (HD, HN, or L) and were declared obsolete in 1971. Animals were dosed with "test article" [either neat HD, 10% agent/chloroform solutions or product solutions (wastestreams) and evaluated for skin-damaging effects (gross and light microscopic). Product solutions from the chemical neutralization of CAIS (either agent/chloroform or agent/ charcoal) produced no microblisters. Dermal application of product solutions from the neutralization of neat HD resulted in microvesicle formation (vesication). All agent-dosed (either agent/chloroform solutions or HD) sites exhibited microblisters as well as other histopathologic lesions. Skin irritant effects were consistent with the skin-injurious activity (erythema and edema) associated with the neutralizing reagent [1,3-dichloro-5,5-dimethylhydantoin (DCDMH)].

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Phase Inspected	Inspection Date	Dated Reported to Study Director	Date of Report to Management
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Sample collection	2/20/96	3/4/96	3/4/96
Histology processing	2/22/96	3/4/96	3/4/96
Test system preparation	6/26/96	7/1/96	7 /1/96
Test article administration - dermal	6/26/96	7 /1/96	7/1/96
Histology processing	6/27/96	7 /1/96	7/1/96
Necropsy/tissue collection	6/27/96	7/1/96	7/1/96
Euthanasia	6/27/96	7 /1/96	7/1/96
Histology processing	6/28/96	7 /1/96	7/1/96
Audit study file	8 /9/96	8/9/96	9/16/96
Audit study file	10/8/96	10/8/96	12/5/96
Audit study file	10/15/96	10/15/96	11/14/96
Audit study file	1/7/97	1/7/97	2/28/97
Audit study file	4/4/97	4/4/97	4/25/97
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GLP COMPLIANCE STATEMENT

The percutaneous dosing of hairless guinea pigs with wastestreams, neutralizing solution and known vesicants, and the gross and histopathologic evaluations of skin lesions in this study were performed by Battelle in compliance with the Environmental Protection Agency's (EPA) Good Laboratory Practice (GLP) Standards (40 CFR Part 792). Likewise, evaluation of the analytical method for HD, HN-1 and L in wastestreams and the determination of HD or HD, HN-1 and L concentrations, as appropriate, in wastestreams was accomplished at Battelle in compliance with EPA GLP Standards. Reports on findings from searches of the literature on HD, HN-1 and L degradation and degradation products and their vesicancy potential as well as analyses of wastestreams for degradation products and residual agent concentrations performed elsewhere than the MREF are excepted from this Good Laboratory Practices Compliance Statement. This study was conducted according to the study protocol, as amended, and Battelle's standard operating procedures. Deviations from the protocol or standard operating procedures are documented in Appendix A. The data presented accurately reflect the results of this study.

Carl T. Olson, D.V.M., Ph.D.

Study Director

6/5/97

Date

QUALITY ASSURANCE

The analytical data supplied by the U. S. Army Edgewood Research, Development and Engineering Center (ERDEC) in support of this task were generated under the auspices of the Research and Technology Directorate Quality Assurance Program Plan. Accordingly, the data are supported by written methodology, sample identification records, and suitable instrument maintenance and calibration. The data and supporting records are retained by ERDEC.

DENNIS W. JOHNSON

Quality Assurance Coordinator

Research and Technology Directorate

PREFACE

The work described in this report was authorized by the Project Manager Non-stockpile Chemical Materiel. This work was started in November 1995 and completed in August 1996.

In conducting the research described in this report, the investigators adhered to the "Guide for the Care and Use of Laboratory Animals," National Institute of Health Publication 85-23, 1985, as promulgated by the Committee on Revision of the Guide for Laboratory Animal Facilities and Care of the Institute of Laboratory Animal Resources, Commission of Life Sciences, National Research Council (Washington, DC). These investigations were also performed in accordance with the requirements of AR 70-18, "Laboratory Animals, Procurement, Transportation, Use, Care, and Public Affairs," and the Laboratory Animal Use and Review Committee (LAURC), U.S. Army Edgewood Research, Development and Engineering Center (ERDEC), which oversees the use of laboratory animals by reviewing for approval all ERDEC research protocols requiring laboratory animals. This project, assigned LAURC Protocol No. 21095000A302C, was approved on 5 Dec 1995.

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TABLE OF CONTENTS

1.	INTROD	UCTION	• • •	• • • •	• • • •	• • •	• • •	• • •	• •	• • •	• •	• • •			•	• •	•	11
2. 1	MATERI	ALS AN	D ME	THOD	s		• • •		• •	• • •	٠.	• • •	• •	• •	•	• •	•	12
2.1 2.1.1 2.1.2		micals Agents Chemic	al A	 gent	 Ide	 nti	fic	 cat	io	 n S	et	 S	• •	• •	•	• •	•	12 12
2.1.3		(Synth Neutra														• •	•	12
2.1.4		Neutra Sets (Neutra cess C	Wast liza	estr tion	eams Sol	uti	on.	• • •	• • •	• • •	••	• • •	• • •	• • •	•			12 13
2.3 2.3.1 2.3.2 2.4 2.5 2.6 2.7 2.8	Tec Ana Ani Exp Ani His	hnolog lytica GC-MS NMR Sp mals a erimen mal Pr topath a Anal	y) . l Me Spec ectro nd Ho tal l epara olog:	thod tros osco ousi Desi atio	olog copy py ng gn n an	ries	osi	ing			• • • • • • • • • • • • • • • • • • • •					• • •	•	14 14 15 15 16 17 19 22
3. 1	RESULT	s							• • •							• •	•	23
3.1 3.2 3.2.1 3.2.2 3.3 3.3.1 3.3.2	Der Dat	mistry mal Ef Gross Histop a Anal Gross Histop	fects Patho atho ysis Patho	ologi logi Res	ic F c Fi ults y (E	ind ndi	ing ngs	gs s 	and	d E	der	na)	•••	• • • • • • • • • • • • • • • • • • • •		• •	•	23 27 27 31 40 40 43
4. I	DISCUS	sion	• • • •								• •				•		•	46
5. (CONCLU	SIONS	• • • •												•			53
1	REFERE	NCES	• • • •												•		•	55
1	APPEND	IXES																
	Α.	Study	Prot	toco:	1						• • •						•	60
	В.	Analy	tical	L Me	thod	olo	дУ			• • •					• •			104
	С.	Studie	es Pe	erfo	rmed	at	EF	RDE	c.		• • •		٠.				•	164
	D.	Gross	Lesi	ion <i>i</i>	Appe	ara	nce	e (24-	-hr)	. 			• •			168
	E.	Dosage	e Sit	te Co	ode	and	Hi	.st	opa	th	010	эду					•	204

TABLES

1	Oxidizer/Solvent System Stoichiometry Utilized in the Modified "Blue", "Red", and "Charcoal" Process Chemistries	14
2	Synopsis of Phase II and Phase III Testing Procedures	18
3	Definitions Used in Histopathologic Evaluations and an Explanation of the Grading of Lesion Severity	21
4	Definition of Degrees of Severity used for Histopathologic Evaluation of Vesication (Microblister Formation)	22
5	Detection and Quantitation Limits for GC/MS Analyses	24
6	Comparison of Agent Residue Levels, Major Products/ By-products, and Unknowns in "Archived" Wastestreams Generated from the Chemical Neutralization of CAIS	26
7	Phase II. Skin Reaction (Erythema and Edema) Follow-ing Exposure to HD, Agent/CHCl ₃ Solutions, and Oxidant/Solvent Solution	28
8	Phase III. Skin Reaction (Erythema and Edema) Following Exposure to HD, Agent/CHCl ₃ Solution, or CAIS Wastestreams	29
9	Phase III. Summary Statistics for Erythema, Edema and Lesion Area after Dosing of "Charcoal" Wastestreams	30
10	Phase II. Vesication (Microblister Formation) in Hairless Guinea-Pigs following Dermal Exposure to HD, Agent/CHCl ₃ Solutions, or Neutralizing Solution (DCDMH/CHCl ₃ /t-BuOH)	32
11	Phase II. Summary of Histopathology Results	33
12	Phase III. Vesication (Microblister Formation) in Hairless Guinea Pigs following Exposure to "Archived" RRS Wastestreams, Agent/CHCl ₃ Solutions, or Neat Sulfur Mustard (HD)	35
13	Phase III. Microblister Formation in Hairless Guinea Pigs Following Exposure to Equal Volumes of "Archived" RRS Wastestreams or Agent/CHCl ₂ Solutions	36

14	Phase III. Microblister Formation in Hairless Guinea Pigs following Exposure to "Fresh" RRS Wastestreams, Agent/CHCl ₃ Solutions, or Neat HD	37
	J	
15	Phase III. Summary of Histopathology Results	38
16	Phase III. Summary of Intermediate to Severe Histopathology Results	39
17	Phase III. Summary of Histopathology following Dosing of "Charcoal" Wastestreams	45
18	Phase III. Summary of Intermediate to Severe Histopathology following Dosing of "Charcoal" Wastestream	45
19	Synopsis of Dermal Toxicity Data for CAIS Agents, Agent Degradation Products, RRS Oxidants and Solvents	49
20	Vesication Potential of Various Analogs/Derivatives of Sulfur Mustard	52
	FIGURES	
1	Guinea Pig Skin Exposure Sites	20
2 - a	Typical Microblister in a Hairless Guinea Pig 24 Hours after Exposure to Vesicant	41
2-b	Normal Skin from a Hairless Guinea Pig	42

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EVALUATION OF THE VESICATING PROPERTIES OF NEUTRALIZED CHEMICAL AGENT IDENTIFICATION SET (CAIS) COMPONENTS

1. Introduction

The U.S. Army Project Manager for Nonstockpile Chemical Materiel has a requirement to develop and field an on site system [Rapid Response System (RRS)]¹ to demilitarize the chemical vesicating (blistering) agents sulfur mustard (HD), nitrogen mustard (HN), and Lewisite (L) contained in Chemical Agent Identification Sets (CAIS). Within the CAIS are glass ampules containing: HD (5 percent), HN (10 percent), or L (5 percent) in chloroform; HD, HN, or L in a charcoal matrix; and neat HD. The proposed RRS operation consists of removing ampules from CAIS and crushing them in a decontamination solution under engineering controls. After chemical neutralization of the agents, the wastestreams are to be turned over to a contractor for ultimate disposal by incineration. The intent of the neutralization process is to produce wastestreams that can be handled in a manner similar to that for industrial wastes. The wastestreams are to be regarded as hazardous material and dealt with in accordance with regulatory guidelines.

Previous research efforts related to the demilitarization/detoxification of CAIS² were conducted in the mid 1970s (Rescigno and Duggan, 1977; Rosenberg, 1977), and recent studies were conducted to ascertain the dermal toxicity characteristics of various RRS wastestreams (Olajos, *et al.* 1996). The current studies stemmed from the need to develop effective chemical neutralization processes which assure the reduction of agent vesicancy. This report provides an account of studies on the vesicancy potential of RRS wastestreams generated from the treatment of CAIS with chemical neutralization reagent.

¹RRS is a transportable system for identification, segregation, repackaging and/or treatment of CAIS.

²Chemical detoxification is defined as a process to convert chemical agents to products that do not exhibit the toxic properties of chemical warfare materiel (CWM). This process is also known as chemical neutralization as defined in Army Regulation 385-61.

2. Materials and Methods

2.1 Chemicals

2.1.1 Agents

Sulfur mustard [2,2'-dichlorodiethyl sulfide (HD), CAS #505-60-2] furnished from Medical Research and Evaluation Facility (MREF) stocks was used neat (undiluted) as a positive control article for vesication.³ Lewisite [dichloro-2-chlorovinyl arsine (L)] CAS #541-25-3 was also furnished from MREF stock. U.S. Army Edgewood Research, Development and Engineering Center (ERDEC) provided a 20 percent solution of nitrogen mustard [bis (2-chloroethyl) ethylamine (HN-1), CAS #538-07-8] in chloroform. Ten percent solutions of HD, HN-1, and L in chloroform were prepared within the MREF laboratory and were used, along with neat HD, as control articles to demonstrate the ability of known vesicants to produce microvesicles and other histopathology when used to dose hairless guinea pigs percutaneously.

2.1.2 Chemical Agent Identification Sets (Synthesized)

Actual ampules from CAIS kits were not used; however "CAIS components" were prepared by ERDEC from agent stocks to contain 10 percent agent in chloroform (Chatfield, *et al.* 1995). Chemical Agent Standard Analytical Reference Material (CASARM) grade HD CAS# 505-60-2 (97.5 mole %), nitrogen mustard [bis (2-chloroethyl) ethylamine (HN-1)] CAS #538-07-8 (≥97% by weight), and CASARM grade lewisite [dichloro-2-chlorovinyl arsine (L)] CAS #541-25-3 (97.8 % by weight) from stocks maintained by the Operations Directorate, ERDEC were used in the preparation of synthesized CAIS. CASARM for HN-1 is not available.

2.1.3 Neutralized Chemical Agent Identification Sets (Wastestreams)

Wastestreams were provided by ERDEC, Aberdeen Proving Ground, MD. Wastestreams from the chemical neutralization of "CAIS components" prepared from agent stocks were tested for vesicancy potential. These wastestreams were prepared by ERDEC as follows:

- Wastestreams from the neutralization of neat HD with 1,3-dichloro-5,5-

³The chemical agents found in CAIS include sulfur mustard, nitrogen mustard, or lewisite. Sulfur mustard was used as representative vesicant for these blistering agents.

dimethylhydantoin (DCDMH) in CHCl₃/t-BuOH/3% H₂O. ("Blue" process)

- Wastestreams from the neutralization of 10% HD, HN, or L (agent in CHCl₃) with DCDMH in CHCl₃/t-BuOH/3% H₂O. ("Red" process)
- Wastestreams from neutralization of HD, HN, or L (agent on charcoal) with DCDMH in CHCl₃ (HD, HN samples) and with DCDMH in CHCl₃/t-BuOH/3% H₂O (L sample). ("Charcoal" process)

Two wastestreams ("archived" and "fresh") were prepared for each process - "Blue", "Red", and "Charcoal" - and samples sent to the MREF for analysis of agent content and for vesicancy testing. The stability of the wastestreams under conditions of administration were not determined by MREF personnel. Test articles are "archived" and "fresh" "Blue", "Red", and "Charcoal" wastestreams.

2.1.4 Neutralization Solution

Neutralizing solution was prepared at the MREF to determine the effect on the skin of dosing this solution alone. For testing vesicating potential, a 0.555M 1,3-dichloro-5,5-dimethylhydantoin (FW 197.02) control article neutralizing solution was prepared by adding 10.9 g DCDMH to a 50:50 tertiary butanol:chloroform with 3 percent water solution in a 100-mL volumetric flask and adding sufficient volume of the butanol/chloroform/water solution to bring the volume to the 100-mL mark. DCDMH (CAS #118-52-5) was purchased from Aldrich Chemical Company (St. Louis, MO). Chloroform (CAS #67-66-3; GC/Spectro grade) was purchased from Burdick and Jackson (Muskegon, MI), and tertiary-butyl alcohol (CAS #75-65-0; ACS Reagent grade) from J.T. Baker (Phillipsburg, NJ). Distilled water was further purified using a Millipore (Bedford, MA) reverse osmosis system.

⁴"Archived" "Blue" and "Red" wastestreams were initially analyzed at ERDEC Oct 95 and re-analyzed for agent residual at the MREF and tested for vesicancy. "Charcoal" wastestream initially analyzed at ERDEC Nov 95 was re-analyzed and tested for vesicancy at the MREF.

⁵"Fresh" wastestreams were prepared and initially analyzed at ERDEC and re-analyzed and tested for vesicancy at the MREF.

2.2 Process Chemistry (Chemical Neutralization Technology)

Process chemistry development was dictated by the requirement for both chemical neutralization and effective detoxification of the agent. Formulations of treatment reagent/solvent systems for the chemical neutralization of CAIS, as reported by ERDEC, are presented in Table 1. Using the neutralization processes, the chemical agents in CAIS may undergo oxidation/chlorination/substitution to yield a mixture of products/by-products, and residual agent may also be present in the wastestreams (Olajos, *et al.* 1996).

TABLE 1. OXIDIZER/SOLVENT SYSTEM STOICHIOMETRY UTILIZED IN THE MODIFIED "BLUE", "RED", AND "CHARCOAL" PROCESS CHEMISTRIES

- 1 volume of neat HD treated with 20 volumes of 0.555M 1,3-dichloro-5,5-dimethylhydantoin (DCDMH) in CHCl₃/t-butanol (50/50) with 3% water by volume. ("Blue" Process)
- 1 volume of each 10% HD in CHCl₃, 10% HN in CHCl₃, and 10% L in CHCl₃ treated with 4 volumes of 0.555M 1,3-dichloro-5,5-dimethylhydantoin (DCDMH) in 50/50 CHCl₃/t-butanol with 3% water by volume. ("Red" Process)
- 45% by weight HD and HN-1 on charcoal treated with excess 1,3-dichloro-5,5-dimethylhydantoin in CHCl₃ combined with 43% by weight L with excess 1,3-dichloro-5,5-dimethylhydantoin in CHCl₃/t-butanol (50/50) with 3% water by volume. ("Charcoal" Process)

2.3 Analytical Methodologies

2.3.1 GC-MS Spectroscopy

Chemically-treated (neutralized) CAIS were analyzed for agent residue levels using full scanning gc-ms spectroscopy. GC-MS spectroscopy was conducted at ERDEC on all wastestreams provided to the MREF, and confirmatory gc-ms analysis was also performed at the MREF prior to conducting the bioassays.

Instrumentation used in the ERDEC analysis of "archived" wastestreams (non-quenched samples) was a Hewlett-Packard 5989B MS engine with Chemstation Data System. Analysis conducted at both ERDEC and the MREF on "fresh" wastestreams, using quenching and derivatization techniques (Dr. Samuel Lucas of Battelle), utilized a Hewlett-Packard Model

5970B Mass Selective Detector (MSD) with an HP 5890A GC and HP 61034 CMS. For procedural details, the reader is referred to Rosso (1995), as provided in ERDEC-TR-372 (Olajos, *et al.* 1996), and Appendix B of this report. Quantitation was based on internal standardization (internal standard = 1,2,4,5-tetrachlorobenzene). Calibration standards were as follows: HD (purity 97.5 %), HN-1 (purity 96.5 %) and L (purity 97.8 %).

Product identification of the CAIS wastestreams was accomplished using gc/ms spectroscopy (EI and CI modes). These studies were performed by ERDEC chemists per procedures outlined in ERDEC Analytical Chemistry Method (Rosso *et al.* 1995; See also Appendix C). The predominant instrument for component identification via the chemical ionization (CI) mode was a Finnigan 5100 gc/ms. The mass spectrometer was operated in the CI mode with methane as the CI reagent gas at a source pressure of 0.5 Torr. Scan time was one sec per scan, and the scan range was 60 to 450 amu. Procedural details have been reported (Olajos, *et al.* 1996).

2.3.2 NMR Spectroscopy

Nuclear magnetic resonance (nmr) spectroscopy analyses of "fresh" wastestreams were conducted at ERDEC as an adjunct to gc-ms analyses. These analyses were performed using a Varian Fourier Transform (FT) nmr spectrometer operated at 200 MHZ for ¹H observation and at 50 MHZ for ¹³C observation. Quantitative data were obtained by digital integration of peak areas.

2.4 Animals and Housing

A total of 35 male (approximately 200-350 g and 3 to 4 weeks of age upon receipt), euthymic hairless guinea pigs (Cr1:IAF (HA)-hr BR), procured from Charles River Laboratories (Wilmington, MA; animals supplied from Portage, MI facility), were used in this study. In this species, the percutaneous application of a vesicating agent such as HD produces the formation of microblisters or microvesicles, a separation of epidermis from dermis of two or more cell widths due to destruction of basal cells (Marlow, et al. 1990, Mershon, et al. 1990, Braue, et al. 1992, and Yourick, et al. 1992). This is analogous to the changes seen in man (Papirmeister, et al. 1984).

Animals were quarantined and screened for general condition and health status, and were maintained in a program accredited by the Association for the Assessment and Accreditation of Laboratory Animal Care (AAALAC) International. Ear tags were applied to maintain positive identification, and animals were maintained between approximately 64 and 79 degrees F and 40 to 70 percent relative humidity with a 12-hr diurnal light cycle. Food and water were provided ad libitum and animals were housed individually in polycarbonate cages prior to exposure to "test article". Following treatment, animals were housed individually within a chemical fume hood during the 24-hr post-exposure period. Following recovery from anesthesia, animals were given food and water.

2.5 Experimental Design

The task was designed to be accomplished in three phases. The first phase was analytical chemistry. This included sample analysis, which was preceded by an evaluation of the limit of detection, the limit of quantitation, the linearity of response, and the precision, accuracy, and specificity of the methodology. Following evaluation of the analytical method, analyses of wastestreams for residual HD ("Blue") or HD, HN-1, and L ("Red" and "Charcoal") were to be accomplished.

The objective of phase II was to assess the biologic effects of dosing volume and exposure time for CAIS components. Two sets of experiments were conducted in Phase II. In the first set, each animal was dosed percutaneously with various volumes of each 10% agent in chloroform solution and with neat HD to determine a volume and a duration of exposure for each agent preparation that resulted in consistent production of microvesication. Once a dosing volume and a duration of exposure for each agent solution were selected, the second set of experiments was conducted to verify consistent microvesication following dosing of agent solutions and to assess the extent of skin pathology following dosing of the neutralization solution alone. Dosing volume of neutralization solution was based upon the approximate volume used to neutralize that volume of agent/agent solution that was determined to consistently create microvesication. Up to eight sites on each guinea pig were dosed and at approximately 24 hr after dosing, using a modification of the Draize method (Draize, et al. 1944), the extent of erythema and edema was graded and

lesion size was measured at each site. The animals were then sacrificed and skin samples taken from dosed sites and prepared for histopathologic evaluation. Histopathologic lesions (microblisters, epidermal necrosis, follicular necrosis, dermal necrosis, vascular necrosis, hemorrhage, and pustular epidermitis) were graded by a veterinary pathologist. Eleven animals were dermally dosed with 1 μ L neat HD and with 10 percent agent (HD, HN or L) in chloroform solutions with dosing volumes ranging from 5 to 50 μ L and exposure times of 1 or 2 hours. Five of these 11 guinea pigs were treated with the DCDMH oxidant/solvent neutralizing solution.

Phase III was to demonstrate that the neutralization process substantially reduced the vesicating properties of wastestreams. Each animal was dosed percutaneously with agents using parameters established in Phase II and with volumes of wastestreams (25 μ L) selected such that the volume of agent in the "Blue" or "Red" treated wastestream that potentially could not be neutralized was approximately equivalent to that agent quantity which when dosed on animal backs created microvesication. For consistency, the same 25 μ L of "Charcoal" wastestream was dosed. The exposed skin was examined 24 hours after dosing and then harvested for histopathologic examination.

The experimental protocol is attached as Appendix A. A synopsis of the experimental design is given in Table 2.

2.6 Animal Preparation and Dosing

Initially using 6 mg xylazine hydrochloride and 35 mg ketamine hydrochloride per kg body weight given intramuscularly and increasing this to 13 mg xylazine and 87 mg ketamine/kg following the first day of dosing, anesthetized guinea pigs were dosed percutaneously on both sides of the dorsal midline with "test articles" (six to eight exposure sites/animal) - see Fig. 1. Table 2 presents a synopsis of treatments, application volumes, and exposure durations. Techniques for dosing are described in Battelle SOP MREF II-009 (Appendix A). Following the requisite exposure time, the exposed skin was decontaminated with 0.5 percent sodium hypochlorite solution (non-irritating concentration). Approximately 24 hours after dosing, the animals were again anesthetized, sites evaluated for erythema and edema and lesion size, and animals then sacrificed with an inhalation anesthetic (halothane) overdose. Following euthanasia,

TABLE 2. SYNOPSIS OF PHASE II AND PHASE III TESTING PROCEDURES

Phase II. A total of 11 hairless guinea pigs (HGPs) were used in this phase with seven or eight sites dosed on each animal. All animals were examined 24 hr following exposure and sites evaluated for erythema and edema. Following this evaluation, animals were euthanatized and dosed skin harvested for histopathology studies.

		Duration of	Total No. HGPs/Total No.
Test Article	Volume Dosed (μL)	Exposure (hr)	Sites Dosed
10% Agent/Chloroform			
HD, HN, or L	5	2	2/2 for each agent solution
	10	2	4/4 for each agent solution
	50	2	2/2 for each agent solution
	5	1	7/7 for each agent solution
	10	1	2/2 for each agent solution
Neat HD	1	2	4/4
	1	1	7/7
Oxidant/Solvent	20	1	5/20

Phase III. A total of 24 HGPs were used, dosing six to eight sites per animal with a 1 hr duration of exposure. All animals were examined 24 hr following exposure and sites evaluated for erythema and edema. Following this evaluation, animals were euthanatized and dosed skin harvested for histopathology studies. Two separate "wastestreams" from each process were provided for testing. The initially provided wastestreams were labeled "archived" and the second set "fresh".

Test Article	Volume Dosed (μL)	Total No. HGPs/Total No. Sites Dosed
10% Agent/Chloroform		
HD, HN, or L	5	20/20 for each agent solution
HD, HN, or L	10	4/4 for each agent solution
Neat HD	1	16/16
Wastestream	,	
"Archived" "Blue" (11/28/95) ^a	25	8/8
"Archived" "Red" (11/28/95)	25	8/8
"Archived" "Charcoal" (1/25/96)	25	8/8
"Archived" "Blue" (11/28/95)	10	4/4
"Archived" "Red" (11/28/95)	10	4/4
"Archived" "Charcoal" (1/25/96)	10	4/4
"Fresh" "Blue" (6/19/96)	25	8/16
"Fresh" "Red" (6/19/96)	25	8/16
"Fresh" "Charcoal" (8/29/96)	25	4/12

a Date is when wastestream was received at the MREF.

skin samples were collected and processed for histopathology.

2.7 Histopathologic Analysis

Following euthanasia, skin from the dosed sites was taken and placed in buffered formalin. After fixation, embedding, and sectioning, skin samples were stained with hematoxylin and eosin and evaluated for histopathology. Histopathologic lesions (microblisters, epidermal necrosis, follicular necrosis, dermal necrosis, vascular necrosis, hemorrhage, and pustular epidermitis) were graded on a scale of 0-4, where 0 = normal, 1 = minimal, 2 = intermediate, 3 = moderate, and 4 = severe. Definitions for scoring of histopathology and the criteria for grading severity of lesions are summarized in Table 3. The grading of microblister formation is highlighted in Table 4.

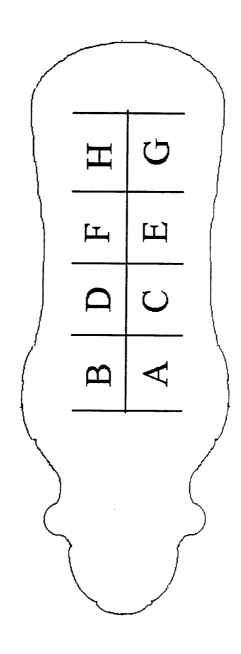


TABLE 3. DEFINITIONS USED IN HISTOPATHOLOGIC EVALUATIONS AND AN EXPLANATION OF THE GRADING OF LESION SEVERITY

30 23/

Microblister	Loss of epidermal basal cell attachment to the underlying basement membrane of at least two adjacent cells. The loss of attachment creates a space which may appear empty, full of proteinaceous fluid, or filled with neutrophils. One or a few isolated small areas of detachment is graded 1, minimal. Many such areas of detachment, or several larger (10 or more contiguous cells) areas of detachment is graded 2, intermediate. When half or more of the epidermis in the tissue section is detached from the dermis, it is graded 3, moderate. Such lesions typically have a much larger space between the basal cells and the dermis. When nearly all of the epidermis is separated from the dermis, it is graded 4, severe. In such situations, there are usually focal, point attachments, so the entire epidermis is not lifted along the full width of the section.
Epidermal necrosis	The epidermal cells exhibit cytoplasmic eosinophilia, nuclear loss or pyknosis, and are generally shrunken. If only individual cells are affected, it is graded 1 (these are generally isolated basal cells). If small areas are affected, with normal areas in close proximity, it is graded 2. If the epidermis exhibits cell death in a full-thickness (all layers of epidermis) pattern, and affects half or more of the skin section, it is graded 3. If the epidermis is virtually entirely necrotic, it is graded 4. Severe ulcers assume that the epidermis is necrotic.
Follicular necrosis	If isolated epithelial cells of the hair follicles exhibit eosinophilia or pyknosis, it is graded 1. If clusters of adjacent cells within follicles are dead, it is graded 2. If cells of half or more of a particular hair follicle are dead, it is graded 3. Grade 4 lesions have complete necrosis of the follicular epithelium underlying much of the epidermal lesion area. This indicates that the agent has penetrated deeply.
Dermal necrosis	Loss of collagen fiber integrity, evidenced by pale eosinophilic staining and homogeneous appearance, indicates necrosis of dermal fibers. With only isolated areas, it is graded 1. Multiple areas are graded 2. Necrosis of most of the superficial dermal collagen in the lesion area is graded 3. A grade 4 lesion requires deep (to the base of the associated adnexa) dermal necrosis.
Vascular necrosis	Loss of integrity of a medium to large blood vessel is vascular necrosis. Grading depends upon the number of vessels affected and the severity. Partial necrosis of one vessel is graded 1 to 2. Complete necrosis of a vessel is graded 3; multiple such lesions are graded 4.
Hemorrhage	Extravasated crythrocytes is hemorrhage. A few isolated foci is graded 1. Multiple, common foci is graded 2. Large pools of blood is graded 3. A grade 4 lesion requires a massive area of blood pooling and the displacement of large areas of dermal collagen.
Pustular epidermitis	Collections of neutrophils in the epidermis proper is graded by extent; one or two small foci is graded 1; three or more small foci is graded 2; one or more large foci is graded 3; a grade 4 lesion would indicate massive infiltration of the entire epidermis by neutrophils.

TABLE 4. DEFINITION OF DEGREES OF SEVERITY USED FOR HISTOPATHOLOGIC EVALUATION OF VESICATION (MICROBLISTER FORMATION*)

Lesion Characteristic	Degree of Severity
No lesion (unaffected)	0 (normal)
One or a few isolated areas of detachment	1 (minimal)
→ Many small areas of detachment or several larger areas of detachment	2 (intermediate)
>50% of the epidermis in tissue section is detached from the dermis (much	3 (moderate)
larger space between basal cells and dermis)	
Nearly all the epidermis is separated from the dermis	4 (severe)

^a Microblister: loss of epidermal basal cell attachment to underlying basement membrane of at least two adjacent cells. Loss of attachment creates a space.

2.8 Data Analysis

For chemistry data generated in Phase I, means and standard deviations of responses of each control standard were determined to calculate both the inter- and intra- variability of the analytical method. Calibration performance characteristics for each analyte, such as slope and standard error of the slope, R² (measure of fit about the regression line), method detection limits, and quantitation limits were calculated.

For Phase III data (vesicating assessment of wastestreams), statistical hypothesis tests were conducted at the 5 percent significance level to determine whether or not the neutralization process reduced the vesicating property of agents contained in CAIS. For each CAIS sample, the incidence of microblisters at sites treated with CAIS agent(s) were compared to those of contralateral sites treated with the wastestream. Although incidence of microblisters was the primary endpoint for evaluating the efficacy of each neutralization process, analyses were also conducted on other indices of skin injury (gross and microscopic). To accommodate the intra-animal correlation of multiple measurements made on the same animal, McNemar's test was used to analyze quantal data (Agresti, 1990). Analysis of variance (ANOVA) models, that include random effects for animal, were fitted to continuous data. If data were not approximately normal, ANOVA were conducted on transformed data, or nonparametric or categorical methods of analysis were performed.

3. Results

3.1 Chemistry

Nitrogen mustard, sulfur mustard, and lewisite are components of CAIS that were chemically neutralized ("detoxified") on reaction with treatment reagent (1,3-dichloro-5, 5-dimethylhydantoin). The selection of a particular process chemistry (designated as "Blue", "Red", or "Charcoal" process) was dependent on whether the agent was neat material (HD), in solution (agent in chloroform), or adsorbed on charcoal.

The DCDMH-mediated neutralization of sulfur mustard resulted in HD concentrations below 50 ppm in "Blue" process wastestream (product solution). The DCDMH reaction resulted in the conversion of sulfur mustard to HD sulfoxide degradation products. Secondary reactions (i.e., elimination/substitution) also occurred which produced chlorinated and vinyl sulfoxides (Olajos, et. al., 1996).

The neutralization reaction between oxidant (DCDMH) and CAIS containing agent (HD, HN or L in chloroform - "Red" Process) resulted in fairly complex product solutions containing various products/by-products and residual amounts of un-reacted agent. Agent levels in "Red" process wastestreams were below 50 ppm for each agent (HD, HN or L) (Ibid).

The process chemistry for the neutralization of CAIS components containing agent (HD, HN or L) on charcoal ("Charcoal" process), resulted in the formation of complex product solutions. Agent residue levels were below 50 ppm for HD and HN and below 85 ppm for L (Ibid).

The wastestreams were complex mixtures which pushed the analytical methodologies to the limits of sensitivity, mixture analysis capability, and structural elucidation. "Archived" and "fresh" wastestreams produced from the neutralization reactions were analyzed by gc-ms for agent residual both at ERDEC and at the MREF. The Method Quantitation Limit (MQL), the concentration level that can be quantitatively reproduced, varied with agent and the matrix analyzed. The analytical capability of the gc-ms method for HD and HN was improved via the utilization of quenched samples (see Table 5). The MQL for all agents in non-quenched samples was $50 \mu g/mL$ (50 ppm) as reported by Rosso (1995) and cited by Olajos, *et al.* (1996). For the quenched samples, the MQL was $15 \mu g/mL$ (15 ppm) for HD or HN and $15 \mu g/mL$ (15 ppm) for

L. Derivatization of lewisite as an enhancement technique in its analysis did not lower the MQL to the desired level (<50 ppm); however, the derivatization prior to quenching did provide reduced variance in the data. See Appendix B for evaluation of the analytical technique and analyses of wastestreams performed at Battelle. It has been proposed that the MQL for L may be lower than 85 µg/mL (Lucas, 1997).

The "archived" wastestreams were additionally analyzed for product/by-product composition at ERDEC (See Table 6 and Appendix C for analytical chemistry performed at ERDEC). HD sulfoxide and other degradation products resultant from secondary reactions (e.g., elimination, substitution) were detected in wastestream samples. HD sulfone and/or its vinyl containing derivatives, which are known vesicants, were not detected in the "Blue" process wastestream. Product analysis did not reveal HD sulfone or vinyl/divinyl analogs in the product solution obtained from the chemical neutralization of the "Red" wastestream. Product characterization of the "Charcoal" wastestream did not reveal HD sulfone; however, multichlorinated vinyl containing derivatives (non-vesicant) were present in the product

TABLE 5. DETECTION AND QUANTITATION LIMITS FOR GC/MS ANALYSES

	Process Wastestreams							
Analyte	"В	lue"	"R	ed"	"Cha	rcoal"		
	MDL ^a	MQLb	MDL	MQL	MDL	MQL		
HD°		50 ppm		50 ppm		50 ppm		
HD ^d	3 ppm	10 ppm	2 ppm	5 ppm	4 ppm	14 ppm		
HN-1°	(*) ^f	(*) ^f		50 ppm		50 ppm		
HN-1 ^d	(*) ^f	(*) ^f	2 ppm	7 ppm	2 ppm	6 ppm		
L°	(*) ^f	(*) ^f		50 ppm		50 ppm		
L ^{d,e}	(*) ^f	(*) ^f	14 ppm	46 ppm	25 ppm	85 ppm		

^{*}MDL = Method Detection Limit

bMQL = Method Quantitation Limit

^{&#}x27;non-quenched sample (Analysis conducted at ERDEC)

dquenched sample (Analysis conducted at Battelle's MREF)

^{*}derivatized lewisite (L-der)

f"Blue" process wastestream contains no HN or L but only neat HD.

solution. Lack of detection of HD sulfone in these wastestreams does not necessarily indicate the absence of this moiety since detection interferences and/or masking by another analyte may have prevented HD sulfone detection.

-1

Nuclear magnetic resonance (nmr) spectroscopy of "fresh" wastestreams conducted at ERDEC did not detect HD in the "Blue" process wastestream and did not detect any HD, HN, or L in the "Red" or "Charcoal" wastestreams. Numerous peaks were evident which were consistent with agent products/by-products (e.g., ClCH₂CHClS(0)CH₂CH₂Cl from HD degradation; ClNHCH₂CH₂Cl from HN degradation; and ClCH = CHAs(OH)₂ from L degradation). Many unidentified product peaks were also detected during nmr analyses.

COMPARISON OF AGENT RESIDUE LEVELS, MAJOR PRODUCTS/BY-PRODUCTS, AND UNKNOWNS IN "ARCHIVED" WASTESTREAMS GENERATED FROM THE CHEMICAL NEUTRALIZATION OF CAIS* TABLE 6.

		Wastestreams	
Component	"Blue" Process	"Red" Process	"Charcoal" Process
	(mod a mom)	(arca /w ppm)	(arca /w ppin)
田	50 ppm	50 ppm	50 ppm
HN	p(*)	50 ppm	50 ppm
_1	p(*)	50 ppm	50 ppm
HD sulfoxide	•••	••••	(-)
HD sulfone	•••	••••	•(-)
sulfones	•(•)	•••	(6.7%)
sulfoxides (multi-chlorinated) ^f	(23.9%)	» (-)	(1.0%)
vinyl derivatives	(0.4%)	(-)	•••
chlorinated alkanes	(0.5%)	(5.3%)	(17.5%)
chlorinated alkenes	(1.1%)	(2.4%)	(40.8%)
other	(25.4%)	(28.4%)	(28.2%)
unknowns	•(•)	(1.8%)	(4.6%)
solvents (CHCl ₃ : t-BuOH)	(48.5%)	(60.4%)	•
Totals	(%8'66)	(98.3%)	(98.8%)

All data derived from gc-ms analysis of sample wastestreams was conducted at ERDEC.

Values based on analysis of non-quenched/non-derivatized samples via gc-ms (CI mode for components analysis; EI mode for agent residual).

Area % calculated from the Total Ion Chromatogram (TIC) of the mass spectrometer. The area % is semi-quantitative; the intent is to show the area under the

peak in comparison to other peaks in the chromatogram.

(*) "Blue" process wastestream contains no HN or L since the CAIS contains only HD.

-) Denotes not detected.

(e.g., tri, tetra-chloro derivatives of HD sulfoxide).

3.2 Dermal Effects

3.2.1 Gross Pathologic Findings

Phase II. All skin exposures to HD and agent/chloroform solutions containing 10 percent HD, HN or L resulted in gross skin lesions consisting of well-defined areas of edema and erythema of moderate to severe intensity. In some instances, large areas of ulceration with complete loss of the covering epidermis was evident. The skin irritant effects of HN and L were comparable to that produced by HD (refer to Table 7 and Appendix D). The skin-injurant effect of oxidant/solvent solution was minimal gross lesions (refer to Table 7 and Appendix D).

Phase III. The cutaneous injury (non-vesicant) effects after 1 hour exposure to HD, agent/CHCl₃ solutions, or CAIS wastestreams ("archived" and "fresh") were evaluated and are summarized in Table 8. Individual gross pathology data are presented in Appendix D. All agent-dosed sites demonstrated gross lesions. Skin lesions were assumed to be elliptical in shape, and lesion area was computed using the formula lesion area = length x width x $\pi/4$. Wastestream-induced dermal injury resulted in mild to moderate degrees of erythema and edema. Because a "fresh" "Charcoal" wastestream was not available for dosing with "fresh" "Red" and "Blue" wastestreams in June, results following dosing of the "fresh" "Charcoal" wastestream in August were combined with results of "archived" "Charcoal" wastestream dosing in March for statistical analyses. Results of these analyses are tabulated in Table 9.

TABLE 7. PHASE II - SKIN REACTION (ERYTHEMA AND EDEMA)
FOLLOWING EXPOSURE TO HD, AGENT/CHCL, SOLUTIONS,
AND OXIDANT/SOLVENT SOLUTION

Experiment Date/ Animal ID	Test Article	Dose Volume (µL)	Time to Decontamination (hr)	No.of Animals Tested	Erythema Score, Mean	Edema Score, Mean
	10% L/CHCl ₃	10	2	2	3.0	3.0
	10% L/CHCl,	50	2	2	3.0	3.0
	10% HN/CHCl3	10	2	2	2.0	2.0
02/19/96 (301, 305)	10% HN/CHCl3	50	2	2	2.0	2.0
(501,505)	10% HD/CHCl ₃	10	2	2	2.0	2.0
	10% HD/CHCl,	50	2	2	2.5	2.0
	Neat HD	1	2	2	2.0	2.0
	10% L/CHCl,	5	2	2	2.5	3.0
	10% L/CHCl ₃	10	2	2	2.5	3.0
00/01/06	10% HN/CHCl ₃	5	2	2	2.0	2.0
02/21/96 (306, 309)	10% HN/CHCl,	10	2 '	2	2.0	2.5
	10% HD/CHCl,	. 5	2	2	3.0	3.0
	10% HD/CHCl,	10	2	2	2.0	2.0
	Neat HD	1	2	2	2.0	2.5
	10% L/CHCl ₃	5	1	2	3.0	3.0
,	10% L/CHCl ₃	10	1	2	3.0	3.0
	10% HN/CHCl3	5	1	2	2.0	2.5
02/27/96 (312, 316)	10% HN/CHCl,	10	1	2	2.0	2.0
(312, 310)	10% HD/CHCl3	5	·1	2	3.0	2.0
	10% HD/CHCl,	10	1	2	2.5	2.0
	Neat HD	1	1	2	3.0	2.5
	10% L/CHCl ₃	5	1	5	3.0	2.8
03/05/96	10% HN/CHCl3	5	I	5	1.8	2.0
(311, 313, 315, 317,	10% HD/CHCl,	5	1	5	2.4	2.4
324)	Neutralizing Solution	20	1	5	0.0	1.0
	Neat HD	1	1	5	2.4	2.6

TABLE 8. PHASE III. SKIN REACTION (ERYTHEMA AND EDEMA) FOLLOWING EXPOSURE TO HD, AGENT/CHCI, SOLUTION OR CAIS WASTESTREAMS

Date,		Dose	No. of	Erythema Score	1 Score	Edema Score	Score	Lesion Area (mm²)	(աա)
Wastestream	Test Article	(nL)	Tested	Mean	S.D.	Mean	S.D.	Mean	S.D.
	10% L/CHCI,	5	8	2.9	0.3	3.0	0.0	95.4	22.2
70,01,00	10% HN/CHCI,	5	8	2.0	6:0	2.0	0.5	60.1	13.6
03/21/96	10% HD/CHCl,	5	8	2.6	0.7	2.1	8.0	107.9	35.8
	"Red" Wastestream	25	8	1.1 ab.c	0.3	1.8	0.5	237.4 des	71.3
"Archived"	"Blue" Wastestream	25	8	1.9 40	8.0	1.6 **	0.5	236.5 ^{de,f}	72.6
Wasicsifeams	"Charcoal" Wastestream	25	8	0.4 1.6.0	0.2	0.4 a,b,c	0.2	132.9	6'56
	Neat HD	1	8	2.8	0.5	2.9	6.0	180.2	53.1
	10% L/CHCl,	5	8	3.0	0.0	3.0	0.0	156.0	67.1
06/20/96,	10% HN/CHCl,	5	8	1.9	0.3	2.1	6.0	82.0	30.8
96/92/90	10% HD/CHCl,	5	8	2.4	0.5	2.3	6.0	94.9	18.4
"Fresh"	"Red" Wastestream	25	8	0.3 ab.c	0.3	0.3 abs	6.0	46.2	52.3
Wastestreams	"Blue" Wastestream	25	8	1.8 4.0	0.5	1.6 1,6,0	6.5	220.6 de.f	42.0
	Neat HD	1	8	2.5	0.5	2.4	6.5	126.8	32.6
	10% L/CHCl,	10	4	3.0	0.0	2.8	6.5	212.6	35.6
08/13/06	10% HN/CHCI,	10	4	2.5	9.0	2.3	6.5	155.1	21.4
06/1700	10% HD/CHCl,	10	4	2.3	1.0	2.3	5.0	178.7	34.9
"Archived"	"Red" Wastestream	10	4	1.1 a,b,c	9.0	0.8 %,6,0	1.0	121.140	41.8
Wastestreams	"Blue" Wastestream	10	4	1.3 4,b,c	0.5	1.0 ".b.c	0.0	142.4	42.3
	"Charcoal" Wastestream	10	4	0.4 a,b,c	0.2	0.0 a,b,c	0.0	69.3ªb.c	23.0
90/00/80	10% L/CHCl,	5	4	3.0	0.0	2.8	0.5	113.9	33.1
06/27/20	10% HN/CHCl,	5	4	1.5	9.0	2.0	8.0	91.3	9.61
"Fresh"	10% HD/CHCl,	5	4	3.0	0.0	3.0	0.00	89.5	26.2
Wastestream	"Charcoal" Wastestream	25	4	0.0 4,6,0	0.0	0.0 a.b.c	0.0	0.04,6.0	0.0

Note: All times to decontamination were 1 hr. a Mean is significantly less than that observed on sites treated with L.

Mean is significantly less than that observed on sites treated with HN. Mean is significantly less than that observed on sites treated with HD. Mean is significantly greater than that observed on sites treated with L. Mean is significantly greater than that observed on sites treated with HN. Mean is significantly greater than that observed on sites treated with HD.

TABLE 9. PHASE III. SUMMARY STATISTICS FOR ERYTHEMA, EDEMA, AND LESION AREA AFTER DOSING OF "CHARCOAL" WASTESTREAMS (#)

Agent/ Compound	Dose Volume (µL)	No. of Animals Tested	Erythema Score Mean S	a Score S.D.	Edema Score Mean	Score S.D.	Lesion Area (mm²) Mean S.D.	a (mm²) S.D.
L (10% solution)	\$	12	2.9	0.3	2.9	0.3	101.6	26.4
HN (10% solution)	\$	12	1.8	0.8	2.0	9.0	70.5	21.4
HD (10% solution)	5	12	2.8	9.0	2.4	8.0	8'101	32.9
"Charcoal" Wastestream	25	12	0.3 ^{b,c,d}	0.3	0.3 b.c.d	0.3	9.88	100.7

Note: All times to decontamination were 1 hr.

Pooled data from the "Charcoal" wastestream received 1/25/96 and dosed on 3/13 and 3/21/96 and wastestream received 8/29/96 and dosed the same day. Volume

of "Charcoal" wastestream dosed was 25 µL at each site.
Mean is significantly less than that observed on sites treated with L.
Mean is significantly less than that observed on sites treated with HN.
Mean is significantly less than that observed on sites treated with HD.

3.2.2 Histopathologic Findings

Phase II. Two hour dermal exposures of animals to neat HD (1 μ L) and to various doses (5 - 50 μ L) of agent/chloroform solutions containing 10 percent HD, HN or L resulted in microblister formation of moderate to severe intensity - refer to Table 10. Incidence of histopathologic changes are summarized in Table 11. In some animals, large areas of ulceration with loss of epidermis prevented the occurrence of microblisters. Individual animal histopathology data are presented in Appendix E. Based on the outcome of the two-hour exposure studies, other guinea pigs were dosed with 5 and 10 μ L volumes of 10 percent agent in chloroform solutions and with neat HD (1 μ L) at an exposure duration of one hour. Microblister formation was evident at all sites, unless occurrence was precluded by development of an ulcer, and ranged in severity from moderate to severe. The application of 5 μ L of 10 percent agent/chloroform solution resulted in microblisters of at least intermediate severity. Refer to Table 10 for incidence/response summary and Appendix E for individual histopathologic findings. The oxidant/solvent system was also evaluated for skin effects. Animals treated with oxidant/solvent solution did not manifest dermal lesions other than minimal inflammatory cell infiltration - refer to Table 11 and Appendix E.

Phase III. Twenty-four animals comprising Phase III of the study were treated with "neutralized" CAIS to ascertain the vesicating potential of chemically degraded CAIS. Incidence/response data related to microvesication are summarized in Tables 12, 13, and 14. A summary of histopathologic changes, including vesication, is presented in Tables 15 and 16. Individual histopathology data appear in Appendix E. Eight animals were dosed with "archived" wastestreams, agent/chloroform solutions, and neat HD. Guinea-pigs dosed with HD and agent/chloroform solutions demonstrated at least minimal microvesication along with consistent, marked epidermal and follicular necrosis. The "Blue" process wastestream ("archived"; 25μ L application) resulted in intermediate to severe microblisters and severe epidermal necrosis at all sites dosed (refer to Tables 12 and 16 and Appendix E). The impression of the pathologist reading the slides was that lesions did not appear to be "basal cell specific", as chemical blistering agents appear to cause, nor did the lesions resulting from application of the "Blue" wastestream penetrate deeply enough to cause severe necrosis in the follicular epithelium. A photomicrograph

TABLE 10. PHASE II. VESICATION (MICROBLISTER FORMATION) IN HAIRLESS GUINEA PIGS FOLLOWING DERMAL EXPOSURE TO HD, AGENT/CHCI, SOLUTIONS, OR NEUTRALIZING SOLUTION (DCDMH/CHCI,/t-BuOH)

				Micro	Microblister Severity (0-4)	0-4)				
Treatment b	Animal									Mean
Group	No.	301	305	306	309				Response	Seventy
Neat HD (1 μ L) 10% HD/CHCI,		2	2	ا و	3				4/4	2.0
, S0 µL		7	7						2/2	2.0
10 µL		7	8	8	8				4/4	8
3μ S				٦,	0				1/2	0.5
10% HN/CHCI,										
20 μL		7	7						2/2	2.0
10 µL		7	7	7	4				4/4	2.5
5 μL				4	4				272	4.0
10% L/CHCI,										
20 ηΓ		4	æ						2/2	3.5
10 hL		ю	ю	m	4				4/4	3.3
5 µL				4	4				2/2	4.0
Treatment d	Animal									Mean
Group	No.	312	316	311	313	315	317	324	Response	Severity
Neat HD (1 μ L) 10% HD/CHCI,		3	3	3	2	2	2	3	חר	2.6
10 μL		М	ю						2/2	3.0
5 μL		6	٣	7	3	່ ຕາ	7	4	L/L	2.9
10% HIN/CHCI,										
10 µL		4	က						2/2	3.5
5 μL		ю	4	æ	4	7	e	4	רור	3.3
10% L/CHCI,										
10 µL		က	4						2/2	3.5
5 μL		m	4	æ.	4	4	7	4	רור	3.4
DCDMH/CHCI,/				0	0	0	0	0	0/5	0
t-BuOH (20 .d.)										

a At 24 hr after dosing, animals were evaluated for skin injury, sacrificed, and skin samples taken and prepared for histopathology.
 b Exposure duration 2 hr.
 c Presence of ulcer prevented formation of microblister.
 d Exposure duration 1 hr.

TABLE 11. PHASE II. SUMMARY OF HISTOPATHOLOGY RESULTS

Animal ID Article 10% L/CHCl, 10% L/CHCl, 10% HN/CHCl, 10% HN/CHCl, 10% HD/CHCl, 10% HD/CHCl, 10% HD/CHCl, 10% HD/CHCl, 10% HD/CHCl, 10% HD/CHCl,		Volume 10 10 50 50 50 50 50 50 50 50 50 50 50 50 50	Decon. 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	No.of Animals 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	No. of Sites 2 2 2 2	Micro- blister 2	Epidermal Necrosis 2	Follicular Necrosis	Pustular Epidermitis	Dermal Necrosis	Hemorrhage	Vascular Necrosis
 	CHCI, CCHCI, VCHCI, VCHCI, VCHCI, CHCI, CHCI,	01 00 20 00 00 00 00 00 00 00 00 00 00 00	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	2 2 2 2 2 2	2 2 2	2	2				- T	
	/CHCI, 4/CHCI, //CHCI, //CHCI, //CHCI, //CHCI,	50 10 50 10 50	2 2 2 2 2	2 2 2 2 2 2	2	4	2	2	0	0	2	0
	V/CHCI, V/CHCI, V/CHCI, V/CHCI, LHD	10 50 10 50	2 2 2	2 2 2 2	2	2		2	0	0	_	0
	VCHCI, VCHCI, VCHCI, IHD	50 10 50	2 2 2	2 2 2		2	2	2	0	0	0	0
	O/CHCI, D/CHCI, tHD	10	2	2	2	2	2	2	0	0	0	0
10% HD Neat	уснсі,	50	2	2	2	2	2	2	0	0		0
Neat	(HD			-	2	2	2	2	0	0	1	0
1/1%01		1	2	2	2	2	2	2	0	0	0	0
10.01	10% L/CHCI,	5	2	2	2	2	2	2	0	1	1	0
10% L/CHCI,	/CHCl,	10	2	2	2	2	2	2	0	1	1	0
NH %01	10% HN/CHCl,	5	2	2	2	2	2	2	1	0	0	0
	10% HN/CHCI,	10	2	2	2	2	2	2	1	1	0	0
(306, 309) 10% HD/CHCI,	женсі,	5	7	2	2	-	2	2	0	2	0	0
10% HD/CHCl,	CHCI,	10	2	2	2	2	2	2	1	1	0	0
Neat HD	t HD	-	2	2	2	2	2	2	0	1	0	0

TABLE 11. PHASE II. SUMMARY OF HISTOPATHOLOGY RESULTS (CONT'D)

Experiment			Time to					Numi	Number of Animals with Sign	th Sign		
Date/ Animal ID	Test Article	Volume (µL)	Decon. (Hr)	No. Of Animals	No. of Sites	Micro- blister	Epidermal Necrosis	Follicular Necrosis	Pustular Epidermitis	Dermal Necrosis	Нетоптивое	Vascular
	10% L/CHCl3	5	1	2	2	2	2	2	0	0	2	0
	10% L/CHCI,	10	1	2	2	2	2	2	0	0	2	0
02/27/96	10% HN/CHCI,	5	1	2	2	2	2	2	2	0	0	0
9	10% HN/CHCI,	10	1	2	2	2	2	2	2	0	0	0
(312, 316)	10% HD/CHCI,	5	-	2	2	2	2	2	_	-	0	0
	10% HD/CHCI,	10	-	2	2	2	2	2	_	0	-	0
	Neat HD		1	2	2	2	2	2	1	0	0	0
	10% L/CHCl ₃	5	1	5	5	5	5	\$	0	1	5	0
96/50/60	10% HN/CHCI,	5		5	5	5	5	\$	4	_	_	0
(311, 313,	10% HD/CHCI,	5	-	5	5	5	5	5	1	3	2	0
315,317, 324)	Neutralizing Solution	20	-	\$	20	0	0	0	0	0	0	0
	Neat HD		-	5	5	5	5	5	0	0	-	0

TABLE 12. PHASE III. VESICATION (MICROBLISTER FORMATION) IN HAIRLESS GUINEA PIGS FOLLOWING EXPOSURE TO "ARCHIVED" RRS WASTESTREAMS, AGENT/CHCI, SOLUTIONS, OR NEAT SULFUR MUSTARD (HD) 4,b

-**----**:

				•		Microblis	Microblister Severity (0-4)	(0-4)			
Treatment Group	Animal No.	494	496	497	499	310	491	493	498	Response	Mean Severity Score
Neat HD (1 μ L)		3	2	7	ε	-	7	2	-	8/8	2.0
10% HD/CHCI,	(5 µL)	-	0	7	4	7			æ	8/L	1.8
10% HN/CHCI,	(5 µL)	8	0		4	4	4	4	7	8/L	2.8
10% L/CHCI,	(5 µL)	4	1	4	3		4	7	æ	8/8	2.8
"Blue" wastestream	(25 µL)	2	4	2	2	2	ю	4	ъ	8/8	2.8
"Red" wastestream	(25 µL)	0	0	0	0	0	0	0	0	8/0	0
"Charcoal" wastestream	(25 µL)	0	0	0	0	0	0	0	0	8/0	0

a Each animal was dosed percutaneously (1 hr exposure) with neat HD, agent/ CHCl, solution, and "archived" wastestreams. Sites were evaluated visually at about 24 hr after dosing, and the animals then sacrificed and skin samples taken and prepared for histopathologic evaluation.

b Dosing volumes of HD and agent/CHCl, solutions, as well as the duration of exposure, were based on preliminary tests. Dosing volumes of wastestreams were based upon approximate ratio of neutralization solution volume to volume of agent treated.

c Wastestreams were generated from the reaction of DCDMH (oxidant) with neat HD ("Blue" process), with 10% HD, HN or L in CHCl, ("Red" process), or with HD, HN, or L on charcoal ("Charcoal" process).

TABLE 13. PHASE III. MICROBLISTER FORMATION IN HAIRLESS GUINEA PIGS FOLLOWING EXPOSURE TO EQUAL VOLUMES OF "ARCHIVED" RRS WASTESTREAMS OR AGENT/CHCI, SOLUTIONS A

Microblister Severity (0-4)

Treatment		- Animal			000	Ş	Description	Mean Severity Score
Group		No.	383	385	389	400	Neshouse	
10% HD/CHCl,	(10 hL)		7	ю	8	ю	4/4	2.5
10% HN/CHCI,	(10 µL)		ю	4	ю	4	4/4	3.5
10% L/CI1Cl,	$(10~\mu L)$		3	4	2	ю	4/4	3.0
Blue" wastestream	(10 hL)		0	şi.m	7	3	3/4	1.5
"Red" wastestream	(10 hL)		0	0	0	0	0/4	0
b "Charcoal" wastestream	(10 pd.)		0	0	0	0	0/4	0

a Each animal was dosed dermally (1 hr exposure) with agent/CHCI, solutions and wastestreams. Sites were evaluated visually at about 24 hr after dosing, and the animals then sacrificed and skin samples taken and prepared for histolopathologic evaluation.

b Wastestreams (product solutions) generated from reaction of oxidant (DCDMH) with HD - "Blue"; 10% HD, HN or L in CHCl3 - "Red"; HD, HN or L on charcoal - "Charcoal".

TABLE 14. PHASE III. MICROBLISTER FORMATION IN HAIRLESS GUINEA PIGS FOLLOWING EXPOSURE TO "FRESH" RRS WASTESTREAMS, AGENT/CHCI, SOLUTIONS, OR NEAT HD ",b"

. (i)

					'	-	Microblis	Microblister Severity (0-4)	ity (0-4)			
Treatment		Animal										Mean Severity
Group	<u> </u>	No.	339	341	342	346	340	345	351	352	Response	Score
Neat HD	(1 µL)		7	7	ю	2	0	-		7	2/8	1.6
10% HD/CHCl,	(5 µL)		ю	7	æ	7	7	7	_	-	8/8	2.0
10% HN/CHCl,	(5 µL)		ю	7	4	4	3	1	-	7	8/8	2.5
10% L/CHCl ₃	(5 µL)		4	ю	ю	7	3	æ	4	æ	8/8	3.1
"Blue" wastestream	(25 µL)		2.5	-	7	-	æ	1.5	7	1.5	8/8	1.8
"Red" wastestream	$(25 \mu L)$		0	0	0.5 ^d	0	0	0	0	0	1/8	0
Treatment Group		Animal No.	379	380	387	388					Response	Mean Severity Score
10% HD/CHCI,	(5 µL)		7	3	3	4					4/4	3.0
10% HN/CHCI3	(5 µL)		33	4	7	33					4/4	3.0
10% L/CHCI,	(5 µL)		4	4	4	4					4/4	4.0
"Charcoal" wastestream	(25 µL)		0	0	0	0					0/4	0

a Each animal was exposed dermally for 1 hr to "test article" (neat IID and/or agent/CHCl, solution, and wastestreams). At 24 hr after dosing, animals were evaluated for gross skin injury and then sacrificed and skin samples taken and prepared for histopathologic evaluation.

b Dosing volumes and duration of exposure were determined from preliminary testing. Dosing volume of wastestreams was selected on the basis of approximate neutralization solution volume to volume to volume of agent. Wastestreams were generated via the reaction of DCDMH with neat HD - "Blue", HD, HN, or L in CHCl, -"Red", and HD, HN, or L on charcoal -

c Mean value for the two sites dosed with each wastestream on each animal.

d Could be due to adjacent HD-treated site.

e Three sites on each animal were dosed with "Charcoal" wastestream and no sites exhibited microblisters.

TABLE 15. PHASE III. SUMMARY OF HISTOPATHOLOGY RESULTS

Date		Dose					Number	Number of Animals with Sign	h Sign		
Source of Wastestream	Agent/ Compound*	Volume (µL)	No. Of Animals	No.of Sites	Micro- blister	Epidermal Necrosis	Follicular Necrosis	Pustular Epidermitis	Dermal Necrosis	Hemorrhage	Vascular Necrosis
	7	5	8	8	8	8	8	0	9	\$	0
20,511.50	HN	5	8	80	7.	8	8	3	4	0	0
03/13/96,	ΩI	5	8	8	7.	8	8	3	L	0	0
	"Red" Wastestream	25	8	æ	p'c'q 0	l b.c.d	p's'q 0	2	p'q 0	0	0
"Archived" Wastestreams	"Blue" Wastestream	25	8	8	8	8	p'o'q l	-	ρl	0	0
	"Charcoal" Wastestream	25	8	8	p's'q 0	9	p'5'q 0	2	p'q 0	0	0
	Neat IID	1	8	œ	8	8	8	0	L	-	0
	7	5	8	œ	8	8	8	1	3	\$	0
06/20/96,	NII	5	8	∞	8	8	8	2	\$	2	0
96/92/90	ŒН	5	8	8	8	8	8	2	\$	\$	0
"Fresh"	"Red" Wastestream	25	8	16	p'z'q l	2 b.c.d	l b,c,d	1	0	0	0
Wastestreams	"Blue" Wastestream	25	8	91	8	8	7	2	1	0	0
	Neat HD	-	8	8	7	8	8	0	\$	5	0
	7	10	4	4	4	4	4	0	0	4	1
76/11/80	HIN	10	4	4	4	4	4	4	1	2	0
	HD	10	4	4	4	4	4	1	0	2	0
"Archived"	"Red" Wastestream	10	4	4	0	0	0	0	0	0	0
Wastesifeams	"Blue" Wastestream	10	4	4	3	3	2	1	0	1	0
	Charcoal Wastestream	10	4	4	0	-	0	-	0	0	0
96/6C/80	-1	5	4	4	4	4	4	0	-	4	0
	NII	5	4	4	4	4	4	1	-	1	0
"Fresh"	ŒI	5	4	4	4	4	4	0	1	2	0
W asiesii caiiis	"Charcoal" Wastestream	25	4	12	0	4	4	_	0	0	0

Note: All times to decontamination were 1 hr.

a Marked ulceration at the dosing site on animal number 496 obscured any evidence of microvesication.

b Incidence of sign was significantly less than that for sites dosed with L using McNemar's Test and a significance level of p=0.05. c Incidence of sign was significantly less than that for sites dosed with HN using McNemar's Test and a significance level of p=0.05. d Incidence of sign was significantly less than that for sites dosed with HD using McNemar's Test and a significance level of p=0.05. c Agent (L, HN, HD) at a concentration of 10% in chloroform.

TABLE 16. PHASE III. SUMMARY OF INTERMEDIATE TO SEVERE HISTOPATHOLOGY RESULTS

4		Dogs				Number	of Animals wit	Number of Animals with Sign Rated Intermediate to Severe	ntermediate	to Severe	
Date, Source of Wastestream	Agent/ Compound	Volume (µL)	No. Of Animais	No.of Sites	Micro- blister	Epidermal Necrosis	Follicular Necrosis	Pustular Epidermitis	Dermal Necrosis	Hemorrhage	Vascular Necrosis
		5	8	8	,9	8	8	0	9	2	0
	HN	5	8	8	,9	8	8	0	3	0	0
03/13/96,	EE	5	8	8	4.	8	8	0	7	0	0
0000000	"Red" Wastestream	25	8	8	5°q ()	0 b,c,d	p,c,d	0	0 b,d	0	0
"Archived"	"Blue" Wastestream	25	8	8	8	8	p'c'q	0	0 p.d	0	0
Wastestreams	"Charcoal" Wastestream	25	8	8	9'q O	2 b,c,d	p'c'q 0	0	p'q 0	0	0
	Neat HD		8	8	•9	8	8	0	9	0	0
	Т	5	8	8	8	8	8	0	3	2	0
06/20/96,	NH.	5	∞	80	,9	8	8	0	3	1	0
96/92/90	Œ	\$	∞	80	.9	8	8	0	\$	2	0
"Fresh"	"Red" Wastestream	25	œ	91	p'5'q ()	p'c'q	J b.c.d	0	0	0	0
Wastestreams	"Blue" Wastestream	25	∞.	91	7	8	2 b.c.d	0	-	0	0
	Neat HD	-	8	8	5	8	8	0	5	-	0
	1	01	4	4	4	4	4	0	0	4	0
20,61,00	HN	10	4	4	4	4	4	0	0	0	0
08/13/96	Œ	10	4	4	4	4	4	0	0	_	0
"Archived"	"Red" Wastestream	10	4	4	0	0	0	0	0	0	0
Wastestreams	"Blue" Wastestream	2	4	4	2	2	0	0	0	0	0
	"Charcoal" Wastestream	01	4	4	0	0	0	0	0	0	0
70/00/80	1	5	4	4	4	4	4	0	-	3	0
08/53/30	AH.	5	4	4	4	4	4	0	0	0	0
"Fresh"	E	\$	4	4	4	4	4	0	-	-	0
Wastestream	"Charcoal" Wastestream	25	4	12	0	0	0	0	0	0	0

Note: All times to decontamination were 1 hr.

a Ulceration at some dosing sites may have obscured evidence of microvesication.

b Incidence of sign was significantly less than that for sites dosed with L using McNemar's Test and a significance level of p=0.05.

c Incidence of sign was significantly less than that for sites dosed with HN using McNemar's Test and a significance level of p=0.05.

d Incidence of sign was significantly less than that for sites dosed with HD using McNemar's Test and a significance level of p=0.05.

e Agent (L, HN, HD) at a concentration of 10% in chloroform.

representative of the morphologic changes observed following treatment with a vesicant is shown in Figure 2a, and one demonstrating the appearance of normal hairless guinea pig epidermis is shown in Figure 2b. The morphologic changes seen consist of ballooning degeneration and loss of epidermal basal cell attachment to the underlying basement membrane. Neither "Red" nor "Charcoal" process wastestreams ("archived"; 25μ L application) produced microblisters (Tables 12 and 15). The "Red" process wastestream produced only minimal pustular epidermitis or minimal epidermal necrosis (refer to Table 15 and Appendix E). The "Charcoal" process wastestream ("archived"; 25μ L application) killed some surface epithelial cells (minimal to intermediate epidermal necrosis) but did not penetrate to basal cells - refer to Table 15 and Appendix E. Four guinea pigs were dosed with 10 μ L of "Blue", "Red", and "Charcoal" process wastestreams ("archived") and evaluated for dermal effect. The "Blue" process wastestream induced microblisters whereas the "Red" and "Charcoal" process wastestreams did not elicit microblister formation. The findings are highlighted in Table 13. Histopathology findings are summarized in Tables 15 and 16, and individual histopathology data are presented in Appendix E.

"Fresh" wastestream-induced skin effects were also evaluated. Data on microvesication are presented in Tables 14, 15, and 16, and other histopathologic skin effects data are given in Tables 15 and 16. Individual animal histopathology results are presented in Appendix E. All agent-dosed sites (neat HD and agent/chloroform solutions) and all "Blue" process wastestream sites demonstrated histopathologic lesions including microvesication. In "fresh" "Red" process wastestream-dosed animals, minimal to no lesions were seen on histopathologic examination. One "Red" process wastestream site in one animal demonstrated histopathology, including minimal microvesication; however, this lesion was incompatible with what had been noted previously. The "Charcoal" process wastestream did not produce microblisters and none of the sites demonstrated histopathology graded more than minimal.

3.3 Data Analysis Results

3.3.1 Gross Pathology (Erythema and Edema)

Means and standard deviations were calculated for erythema and edema scores (Phase II and III Studies) and for lesion areas (Phase III Studies). Analysis of variance was performed for

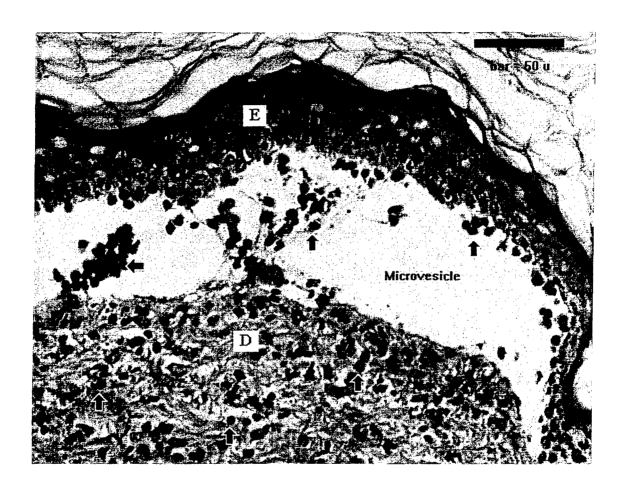


Fig. 2a. Typical microblister in a hairless guinea pig 24 hours after exposure to vesicant. Epidermis (E) is eosinophilic and shrunken due to necrotic epithelium; dermis (D) is also necrotic and contains an infiltrate of polymorphonuclear cells (arrows), as does the microblister cavity (microvesicle).

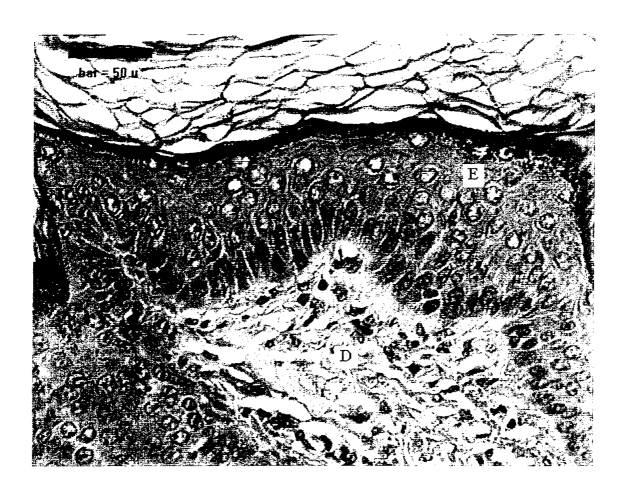


Fig. 2b. Normal skin from a hairless guinea pig. Epidermis (E) and dermis (D) are visible. Note differences in appearance from the necrotic tissue depicted in Fig. 2a. Magnification of both skin photomicrographs is the same.

inflammation scores and lesion areas. Tables 8 and 9 present means and standard deviations for erythema and edema scores. Significant decreases in average inflammation scores resulted when comparing wastestream-dosed ("archived" or "fresh" -25 μ L volume application) to agent-dosed sites (HD or agent/chloroform) - refer to Table 9. Some significant increases in lesion areas were noted with wastestreams, presumably due to the larger volume dosed. For the August 13, 1996 experiment (vesicancy assay of "archived" wastestreams), significant decreases in average inflammation scores as well as average lesion areas resulted when comparing wastestream-dosed ("archived" "Red" and "Blue" process wastestreams - $10~\mu$ L volume applications of wastestreams and agent/chloroform solutions) to agent-dosed sites. All observed inflammation scores and lesion areas from the "fresh" "Charcoal" wastestream-dosed sites were zero.

3.3.2 Histopathology

Statistical analysis (McNemar's test) of the histopathology data was performed to ascertain the significance between treatment groups (neat HD, agent/chloroform solutions, and wastestreams) at the 0.05 significance level. Sites dosed with "Red" or "Charcoal" wastestream ("archived", 25 μ L volume application) exhibited a <u>significant decrease</u> in incidence of <u>microblisters</u> compared to those sites dosed with HD or agent/chloroform solutions. Sites dosed with the wastestreams also showed a significant decrease in the incidence of follicular necrosis compared to sites dosed with any of the three agents (HD, HN, or L in chloroform, neat HD). Some significant neutralized wastestream versus agent differences also resulted with respect to incidence of epidermal and dermal necrosis.

Sites dosed with "Red" wastestream ("fresh", 25 μ L volume application) showed a <u>significant decrease</u> in incidence of <u>microblisters</u>, epidermal necrosis, and follicular necrosis compared to that on sites dosed with any of the three agents. Numerical reductions in some pathology from wastestream-dosed sites ("archived", 10 μ L volume application) were observed, although they were not statistically significant due to the smaller number of animals tested.

Statistical analysis of incidence of intermediate to severe histopathologic signs was also performed. Sites dosed with "Red" or "Charcoal" wastestream ("archived", 25 μ L volume application) demonstrated a <u>significant decrease</u> in incidence of microblisters compared to that on sites dosed with L/chloroform and HN/chloroform. A decrease in incidence was also observed

for the "Red" or "Charcoal" wastestream compared to that on sites dosed with HD/chloroform, but were not statistically significant because only four of the eight animals exposed to HD/chloroform had intermediate to severe microblisters. Sites dosed with "Red" or "Charcoal" wastestream ("archived", 25 µL volume application) demonstrated a significant decrease in incidence of epidermal necrosis and follicular necrosis compared to that on sites dosed with any of the three agents. Sites dosed with "Red" wastestream ("fresh", 25 µL volume application) showed a significant decrease in incidence of microblisters, epidermal necrosis, and follicular necrosis compared to that on sites dosed with any of the three agents. Sites dosed with "Blue" wastestream showed a significant decrease in incidence of follicular necrosis compared to that observed on sites dosed with any of the three agents.

Sites dosed with "fresh" "Charcoal" wastestream ($25 \mu L$ volume application) exhibited a numerical reduction in incidence of microblisters, although this was not statistically significant due to the smaller number of animals tested, compared to that observed on sites dosed with any of the three agents. Statistical analyses also were conducted on the pooled "Charcoal" wastestream data ("fresh" and "archived", $25 \mu L$ volume applications- see Table 17). These analyses assumed that the probability of a microblister and other histopathologic endpoints is similar for sites dosed with "archived" and "fresh" "Charcoal" wastestreams. Pooled data for sites dosed with "Charcoal" wastestream showed a significant decrease in incidence of microblisters and follicular necrosis compared to that on sites dosed with any of the three agents. Statistical analyses of incidence of intermediate to severe histopathologic signs (Table 18) were also performed on the pooled "Charcoal" wastestream data ("fresh" or "archived", $25 \mu L$ volume application). Sites dosed with "Charcoal" wastestream showed a significant decrease in incidence of intermediate to severe microblisters, epidermal necrosis, and follicular necrosis compared to that observed on sites dosed with any of the three agents.

PHASE III. SUMMARY OF HISTOPATHOLOGY FOLLOWING DOSING OF "CHARCOAL" WASTESTREAMS TABLE 17.

						Number o	Number of Animals with Histopathology	pathology		
Agent/ Compound ^r	Volume (µL)	No. Of Animals	No. of Sites	Micro- blister	Epidermal Necrosis	Follicular Necrosís	Pustular Epidermitis	Dermal Necrosis	Hemorrhage	Vascular Necrosis
Π	5	12	12	12	12	12	0	7	6	0
NH	5	12	12	11 ۽	12	12	4	\$	-	0
ŒН	5	12	12	119	12	12	3	88	2	0
"Charcoal" Wastestream	25	12	20	0 دوه	10	4 c.de	3	•>0	٥٤	0

Note: All times to decontamination were 1 hr.

Pooled data from the "Charcoal" wastestream received 1/25/96 and dosed on 3/13 and 3/21/96 and wastestream received 8/29/96 and dosed the same day. Volume of "Charcoal" wastestream dosed was 25 µL, at each site,

Marked ulceration at the dosing site on animal #496 may have obscured microvesication. Incidence of pathology was significantly less than that for sites dosed with L based on McNemar's Test at the 0.05 significance level.

Incidence of pathology was significantly less than that for sites dosed with HN based on McNemar's Test at the 0.05 significance level. Incidence of pathology was significantly less than that for sites dosed with HD based on McNemar's Test at the 0.05 significance level.

Agent (L, HN, HD) at a concentration of 10% in chloroform.

PHASE III. SUMMARY OF INTERMEDIATE TO SEVERE HISTOPATHOLOGY FOLLOWING DOSING OF "CHARCOAL" WASTESTREAM ^{*} TABLE 18.

	Doce				Numbe	r of Animals with	Number of Animals with Histopathology Rated Intermediate to Severe	ated Intermed	late to Severe	
Agent/ Compound	Volume (µL)	No. of Animals	No. of Sites	Micro- blister	Epidermal Necrosis	Follicular Necrosis	Pustular Epidermitia	Dermal Necrosis	Hemorrhage	Vascular Necrosis
Т	5	12	12	10	12	12	0	7	5	0
NH	5	12	12	10	12	12	0	3	0	0
ŒН	5	12	12	8	12	12	0	•	1	0
"Charcoal" Wastestream	25	12	20	0 b.c.d	2 b.c.d	0 b.c.d	0	p'q 0	0	0

Note: All times to decontamination were 1 hr.

a Pooled data from the "Charcoal" wastestream received 1/25/96 and dosed 3/13 and 3/21/96 and wastestream received 8/29/96 and dosed the same day,

Incidence of pathology was significantly less than that for sites dosed with L based on McNemar's Test at the 0.05 significance level.

Incidence of pathology was significantly less than that for sites dosed with HN based on McNemar's Test at the 0.05 significance level. Incidence of pathology was significantly less than that for sites dosed with HD based on McNemar's Test at the 0.05 significance level.

Agent (L, HN, HD) at a concentration of 10% in chloroform.

4. Discussion

The intent of the process chemistries was to develop neutralization reactions that achieved destruction of CAIS agents, forming wastestreams with minimal toxic hazards. Achieving the desired objectives represented a formidable challenge since chemical reactions with the agents can result in the formation of reaction products/by-products having vesicant action and/or a high degree of systemic toxicity. Destruction of agents involves complex chemical reactions, which is certainly the case for sulfur mustard. HD destruction is complicated by the presence of sulfur and chlorine in the HD molecule, which in some cases facilitates and in others impedes the chemical degradation of HD. Possible methods suggested for agent destruction have included oxidation and chlorination for HD and oxidation for nitrogen mustard and lewisite. The toxicity of the degradation products resulting from the chemical neutralization of HD, HN, or L is of concern to the toxicology, health, and regulatory communities. The current studies were undertaken to assess the vesicant properties of neutralized CAIS.

Current methods for demilitarizing CAIS are still based largely on chemical neutralization via oxidizing materials although the use of DCDMH as oxidant does provide an alternative degradation pathway via the chlorination of HD. The oxidation of sulfur mustard, as pointed out by Franke (1967), represents one of the most important decontamination reactions for HD. The oxidation of sulfur mustard via various oxidizers (e.g., hydrogen peroxide, hypochloric acid and its salts, potassium permanganate, nitric acid, DCDMH, etc.) yields various compounds whose composition depends on the nature of the oxidant used and the specific reaction conditions. Most easily formed is HD sulfoxide which on oxidation yields HD sulfone - both represent major oxidation products of sulfur mustard.

The oxidation of HD not only alters the skin-damaging properties of HD but the systemic toxicity of sulfur mustard as well. The oxidation of HD is of great interest since sulfoxide formation, on chemical neutralization of HD, can be considered a "detoxification". The "detoxification" of HD via oxidation to the sulfoxide was demonstrated in the 1940's. In contrast, the formation of mustard sulfone, a product of further oxidation, can contribute to an enhanced systemic toxicity and vesicant potential of the product solution/mixture. HD sulfone, having the

S(O)₂ functional group, is highly poisonous and <u>comparable</u> in toxicity to HD.⁶ Research conducted since Philips' review (Philips, 1950) on sulfur mustard pharmacology/toxicology demonstrated that HD sulfone is a highly toxic vesicant.

Certainly, based on the known toxicity characteristics of mustard sulfone, mustard sulfoxide, and their vinyl derivatives, it is crucial that the process chemistries developed for the destruction of CAIS employ oxidants that minimize the formation of HD sulfone and HD analogs having comparable biological activity (systemic toxicity and vesicancy) to that of HD.

A concern regarding the vesication potential of HD degradation products/by-products prompted a review of the toxicology literature pertaining to sulfur mustard products/by-products information which is summarized in Table 19. The reader is referred to a review on the subject matter (Olajos *et al.* 1996).

For purposes of this report, discussion on the relationship between chemical structure and vesication is limited to the thioether molecule. Degradation product(s) of nitrogen mustards have not been implicated as having vesicant potential although this area of research needs to be explored. The principal degradation product of lewisite, namely L oxide, is a potent vesicant. The reader is referred to several papers/reviews on the subject of mustard vesication and toxicology (Bouder, 1940, Anslow and Houck, 1946, Philips, 1950, Aleksandrov, 1969; Franke, 1967, and Henry, 1991) as well as reviews covering the systemic toxicity and pathology of nitrogen mustards (Anslow and Houck, 1946, Renshaw, 1946, Cope *et al.*, 1946, and Graef *et al.*, 1948). The subject of lewisite toxicology and pathology has also been amply covered (Wardell, 1941, Gates *et al.*, 1946; and Goldman and Dacre, 1989).

The vesicant potential of sulfur mustard derivatives (oxidation and chlorination products) has been investigated since the 1920's to modern times. Research has indicated that the strongest vesicant action is exerted by β -halogenated sulfides. The position and degree of chlorination influences the vesicant potential of the thioether molecule. With respect to the site of chlorination, Kirner (1928) and Dawson and Wardell (1930) concluded that compounds having

⁶ HD is easily destroyed by all chlorinating agents (aqueous or anhydrous medium). Under appropriate conditions, the chlorination of HD can proceed to form various polychlorides. In the presence of water, chlorination of HD is altered resulting in the formation as sulfoxides (Aleksandrov, 1969).

the chlorine atom in the beta position were considerably more vesicant that those having chlorine in the alpha or gamma position. The degree of chlorination also influences the vesicant activity of the sulfide molecule and hence the early use of chlorination to degrade HD. Monosubstitution analogs of HD, regardless of position, are less effective vesicants than HD. As previously stated. the introduction of halogen atoms results in decreased toxicity and markedly diminished vesicant action. Research in the 1920s (Mann and Pope, 1922; Peters and Walker, 1923; and Lawson and Dawson, 1927) - summarized by Bouder (1940) - indicated that the higher chlorinated derivatives (e.g., tri-, tetra-, and hexachloro derivatives) of HD (saturated or unsaturated) were non-vesicant. A summary of the vesicant potential of various chlorinated analogs of sulfur mustard are given in Table 20. The demilitarization of CAIS as stated is based on chemical neutralization via oxidizing materials which not only alters the systemic toxicity of HD (as discussed) but the skin damaging properties (irritation, vesication). Fuson et al. (1943) on review of the vesicant activity of sulfur compounds concluded that compounds containing the S(0) group were non-vesicant. Mustard sulfone, containing the S (0)2, functional group is a known vesicant (vesicancy potential 1/7 to 1/5 of HD; Bergmann et al., 1945). The formation of HD sulfone can contribute to an enhanced vesicant potential of the product solution/mixture (wastestream).

SYNOPSIS OF DERMAL TOXICITY DATA FOR CAIS AGENTS, AGENT DEGRADATION PRODUCTS, RRS OXIDANTS AND SOLVENTS* TABLE 19.

Compound	Dermal Toxicity ^b (LD _{ss} /LDLo/TDLo)	References	Skin Effects (Irritation, Vesication) ^b	References
AGENTS HD [bis(2-chloroethyl)sulfide]	LD ₅₀ (40-100 mg/kg)	Anslow & Houck (1946)	Anslow & Houck (1946) Severe irritant/escharotic, severe vesicant	Marshall & Williams (1921); Gates & Moore (1946);
L [dichloro(2-chlorovinyl)arsine]	LD ₅₀ (5-6 mg/kg)	Cameron et al. (1946); Gates et al. (1946)	Severe irritant/escharotic, severe vesicant	Kensnaw (1946) Gates et al. (1946)
HN-1 [bis(2-chloroethyl)ethylamine]	LD _{so} (15-20 mg/kg)	Smith (1943a); Anslow & Houck (1946)	Severe irritant/escharotic, severe vesicant	Cope et al. (1946); Renshaw (1946)
HN-3 [tris(2-chloroethyl)amine]	LD ₅₀ (5-20 mg/kg)	Smith (1943d); Anslow & Houck (1946)	Severe irritant/escharotic, severe vesicant	Cope et al. (1946); Renshaw (1946);
OXIDIZED DERIVATIVES HD sulfoxide) (-)	3 (-)	Irritant, non-vesicant	Marshall & Williams (1921); Lawson & Dawson (1927); Young et al. (1944)
Sulfoxide, 2-chloroethyl vinyl	p(-)	p(-)	Irritant, non-vesicant	Thomson et al. (1945)
Divinyl sulfoxide	(-)	•(-)	Irritant, non-vesicant	Fuson et al. (1943); Young et al. (1944) Thomson et al. (1945)
HD sulfone) (-)	, (-)	Irritant/escharotic, vesicant	Marshall & Williams 1921); Young et al. (1944)

TABLE 19. (Continued)

Compound	Dermal Toxicity ^a (LD _{so} /LDLo/TDLo)	References	Skin Effects (Irritation, Vesication) ^b	References
OXIDIZED DERIVATIVES (Cont.)	(Cont.)			
Sulfone, 2-chloroethyl vinyl	g (-)	g (-)	Irritant/escharotic, vesicant	Young et al. (1944) Thomson et al. (1945)
Divinyl sulfone	LD_{so} (= 20 mg/kg)	Smyth et al. (1962)	Irritant/escharotic, vesicant	Young et al. (1944); Thomson et al. (1945)
HN-1 oxide	4 (-)	(-)	u(-)	u(-)
HN-3 oxide	; ①	, (-)	u(-)	u(-)
Lewisite oxide	₹.	r (•)	Irritant/escharotic, vesicant	Young et al. (1944); Thomson et al. (1945)
2-chlorovinylarsonic acid	(-) _k	(-) k	Irritant, non-vesicant	Young et al. (1944); Thomson et al (1945)
2-chlorovinylarsonous acid	- (-)	- (·)	Irritant, non-vesicant	Cameron et al. (1946)
OXIDIZERS				
рсрмн	LD ₅₀ (>20 g/kg)	EPA 8EHQ0281-0382; EPA 88-8100-228	Severe irritant	EPA 8EHQ0281-0382; EPA #88-8100-173 (cited in RTECS)

TABLE 19. (Continued)

	Dermal Toxicitya		Skin Effects	
Compound	(LD50/LDLo/TDLo)	References	(Irritation, Vesication)	ion) ^b References
SOLVENTS				
Chloroform	LD _{so} (>20 g/kg)	NTIS AD-A062-138 (cited in RTECS)	Mild irritant	Guido and Martins (1988)
t-butyl alcohol	m (-)	m (-)	Mild irritant	Oettel (1936)

Table modified from that originally compiled by Olajos et al., 1996.

Rabbit as animal model unless otherwise indicated. Tests for irritancy based on animal and/or human studies.

Test for vesicant action of agents conducted on human subjects.

Mouse s.c. LD₃₀ (>25 mg/kg) [Anslow and Houck (1946)]

Rat oral (100 mg/kg, mortality 1/1) [Young et al., 1944] Mouse s.c. LD₅₀ (>25 mg/kg) [Anslow and Houck (1946)]. Mouse s.c. LD₅₀ (>25 mg/kg) [Anslow and Houck (1946)]. Acute toxicity undetermined.

Mouse i.p. LD₁₀ (50-100 mg/kg) [Bergmann and Fruton (1943); Stahmann and Bergmann (1946a)].
 Mouse i.p. LD₁₀ (2-5 mg/kg) [Bergmann and Fruton (1943); Stahmann and Bergmann (1946a)].
 Mouse s.c. (mortalities: 2 mg/kg (0/5); 5 mg/kg (5/5); 10 mg/kg (5/5) [Young et al. (1944)].
 Mouse i.p. [mortalities: (1000 mg/kg 10/10; 500 mg/kg 0/10] (Young et al., 1944).
 Reported as hightly toxic, details not given (Cameron et al., 1946).
 Mabbit oral LDLo (4.5 g/kg) [RTECS].
 Young et al., (1944) reported HN2 oxide as non-vesicant; no data for HN1, HN3.

TABLE 20. VESICATION POTENTIAL OF VARIOUS ANALOGS/DERIVATIVES OF SULFUR MUSTARD^a

Analogs/Derivatives (Saturated and Unsaturated)	Vesicant Activity	References b
OXIDIZED DERIVATIVES		
Mustard Sulfone (sulfone, bis(2-chloroethyl)	(POS)	Marshall & Williams (1921), Young et al. (1944)
Sulfone, 2-chloroethyl vinyl	(POS)	Young et al. (1944)
Divinyl Sulfone	(POS)	Young et al. (1944), Thomson et al. (1945)
Mustard Sulfoxide (sulfoxide, bis(2-chloroethyl)	(NEG)	Marshall & Williams (1921) Lawson & Dawson (1927) Fuson et al. (1943) Bergmann et al. (1945)
Divinyl Sulfoxide	(NEG)	Young et al. (1944) Thompson et al. (1945) Bergmann et al. (1945)
β-chloroethyl vinyl sulfoxide	(NEG)	Young et al. (1944)
α , β' , -trichlorodiethyl sulfoxide	(NEG)	Young et al. (1944)
CHLORINATED DERIVATIVES		
bis(α-chloroethyl) sulfide	(NEG)	Peters and Walker 1923) Baldwin <i>et al.</i> (1924) Kirner (1928) Dawson & Wardell (1930)
α,β,β' -trichlorodiethyl sulfide	(NEG)	Mann & Pope (1922) Lawson & Dawson (1927)
$\alpha,\beta,\beta,\beta'$ tetrachlorodiethyl sulfide	(NEG)	Mann & Pope (1922) Lawson & Dawson (1927)
$\alpha,\alpha^1,\beta,\beta'$ tetrachlorodiethyl sulfide	(NEG)	Lawson & Dawson (1927)
$\alpha,\alpha\beta,\beta,\beta'$ hexachlorodiethyl sulfide	(NEG)	Mann & Pope (1922) Lawson & Dawson (1926) Dawson & Wardell (1930)
β -chloroethyl α , β dichlorovinyl sulfide	(NEG)	Lawson & Dawson (1926) Kirner (1928) Dawson & Wardell (1930)
β -chloroethyl α , β , β' trichlorovinyl sulfide	(NEG)	Lawson & Dawson (1926) Kirner (1928) Dawson & Wardell (1930)
β -chloroethyl chlorovinyl sulfide (α and β isomers)	(POS)	Lawson & Dawson (1926) Dawson & Wardell (1930) Fuson <i>et al.</i> (1943)

⁽a) Table from Olajos et al., 1996 (b) citations are primary and/or secondary

The lack of vesicancy following treatment with "Red" and "Charcoal" process wastestreams is indicative of the effectiveness of the neutralization chemistries in destruction of chemical agent concomitant with the minimization of potentially vesicant-inducing products/by-products. Product analyses corroborated the results of the bioassay. The composite agent (HD, HN and L) levels in "archived" and "fresh" "Red" wastestreams and in "archived" and "fresh" "Charcoal" wastestreams did not elicit vesication in the volumes dosed.

Treatment with "Blue" process wastestreams ("archived" and "fresh") resulted in a vesicant response. The bioassay results were unexpected since the agent residual level was 15 ppm or less, a level below that expected to elicit a vesicant response. The most plausible explanation for vesication is that degradation product(s)/by-product(s) were present in the wastestreams and elicited the vesicant response. The association of vesicancy with the "Blue" process wastestreams ("archived" and "fresh") presents concerns regarding (1) unattained reduction in agent characteristics of the "Blue" wastestreams and (2) needed refinement of the analytical techniques for product identification in the wastestreams.

5. Conclusions

Based on the findings of these studies the following conclusions can be made.

- The vesicating properties of the "Blue" wastestream were not significantly reduced from that of the untreated CAIS (neat HD) prior to treatment with neutralization solution.
- The vesicating properties of both "Red" and "Charcoal" wastestreams, in the volumes dosed, were significantly lower than the untreated CAIS agent solutions.

Blank

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APPENDIX A

Study Protocol
Battelle SOP MREF II-009
Deviation Reports/Memos to File

Study Performed by Battelle Memorial Institute Medical Research and Evaluation Facility 505 King Avenue, Building JM-3, Columbus, OH 43201-2693

STUDY TITLE:

Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components

PRINCIPAL INVESTIGATOR:	Cal T Dlan	11/29/95
SCIENTIFIC REVIEW:	Carl T. Olson, DVM, PhD, S	Study Director
	John B. Johnson, DVM, Mana Research and Evaluation Fa	
ATTENDING/CONSULTING VET	TERINARIAN:	
STATISTICAL REVIEW:	Tracy A. Peace, DVM, Study Ruld G. Ment	Veterinarian
	Ronald G. Menton, PhD, Stu- Statistician	dy

CONTRACTING OFFICER'S REPRESENTATIVE:

Richard R. Stotts, LTC, USA, VC

PROTOCOL TITLE: Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components

PRINCIPAL INVESTIGATOR: Carl T. Olson, DVM, PhD

CO-INVESTIGATOR(S):

Study Supervisor: Robyn C. Kiser, B.S.

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Study Veterinarian: Tracy A. Peace, D.V.M.

Study Pathologist: Allen W. Singer, D.V.M.

Study Chemist: Timothy L. Hayes, B.A.

Sponsor: Program Manager for Non-Stockpile Chemical

Materiel (PMNSCM), USACMDA

<u>Sponsor Monitor</u>: LTC Richard R. Stotts, D.V.M., U.S. Army Medical Research Institute of Chemical Defense (USAMRICD)

I. NON-TECHNICAL SYNOPSIS:

The objective of this study is to assess the vesicating potential of wastestreams following neutralization of chemical agents HD, HN, and L contained in Chemical Agent Identification Sets (CAIS). Experiments are conducted with the hairless guinea pig to evaluate the vesicating properties of wastestreams and to develop a data base to address regulatory concerns of worker safety, industrial hygiene, and transportation and disposal of hazardous waste. This study will be conducted following the guidelines of the Environmental Protection Agency (EPA) Good Laboratory Practice (GLP) Standards.

II. BACKGROUND:

A. Background:

The Program Manager for Non-Stockpile Chemical Materiel has a requirement to develop and field a transportable system to neutralize vesicating (blister) agents contained in CAIS. Within these sets are glass ampules containing either 1) neat sulfur mustard (HD), 2) 5-10 percent HD, nitrogen mustard (HN), or Lewisite (L) in chloroform, or 3) 43 percent by weight HD, HN or L in a charcoal matrix. The proposed operation for neutralization of these vesicating agents consists of removing ampules from containers and crushing them in a

neutralization solution under engineering controls. neutralization solution is 0.555 M 1,3-dichloro-5,5dimethylhydantoin in approximately 50:50 chloroform/tbutanol with about 3 percent water. Volume of neutralization solution to volume of ampules is approximately 20:1 for neat HD, 4:1 for agents in chloroform, and 70:1 for agents on the charcoal matrix. After chemical neutralization of the agents, the wastestreams will be turned over to a hazardous waste disposal contractor for ultimate disposal by incineration. The intent is to have wastestreams handled in a manner similar to industrial wastes that are readily transported and destroyed in accordance with regulatory quidelines. In order to be able to handle the wastestreams as normal regulated industrial hazardous waste, it is necessary to demonstrate that the vesicating properties of the chemical agents have been virtually eliminated.

B. Literature Search:

1. Literature Source(s) Searched:

The Current Contents® monthly computerized Life Sciences database is routinely searched for publications on biological effects of Chemical Surety Materiel (CSM) or dilutions. MEDLINE and TOXLINE are likewise routinely searched. On-line searches were conducted which included DTIC and FEDRIP to ascertain if previous studies (vesication) have been conducted on CAIS based on the process chemistries involved and utilization of the hairless guinea pig.

2. Date and Number of Search:

TOXLINE and MEDLINE were searched in July 95 for papers on sulfur mustard. DTIC and FEDRIP were searched in September 1995.

- 3. Key Words of Search: Key words used in searches included: sulfur mustard, hairless guinea pig, chemical agent identification sets (CAIS), CAIS wastestreams and/or components, and/or process chemicals, e.g. 1,3 dichloro-5,5 dimethylhydantoin (DCDMH), effects (vesication), sulfides, and sulfoxides.
- 4. Results of Searches: The hairless guinea pig model to measure microvesication as an indicator of dermal exposure to "blister agents" is the only well-accepted, published method to assess vesicating potential. No reports on vesication studies on CAIS/CAIS wastestreams in hairless guinea pigs were found.

III. OBJECTIVE/HYPOTHESIS:

The hypothesis of the study is that mean levels of skin irritation endpoints, including microvesication, at sites treated with neutralized CAIS compounds are statistically less than mean levels at sites treated with CAIS compound mixtures.

IV. MILITARY RELEVANCE:

The military relevance is discussed under background.

V. MATERIALS AND METHODS:

A. Experimental Design and General Procedures:

Phase I: Chemical Analyses and Validation of Methods

In the first phase of this task, analytical chemistry methods (e.g., gas chromatography-mass spectroscopy) proposed by ERDEC for determination of concentrations of HD, HN, and L in wastestreams will be evaluated. Modifications to these methods may be made based on the analytical equipment available in MREF laboratories and previous experience of chemists in the analysis of CSM. The need for extensive modifications is not anticipated. Each of the analytical methods, with any modifications, will be validated by evaluating the limit of detection, the limit of quantification, the linearity of response, and the precision, accuracy, and specificity. Analyses of the actual wastestreams for HD, HN and L concentrations will be accomplished following validation of the methods.

Phase II: Dosing Parameters for Agent(s) and Neutralization Solution

The objective of this phase is to assess the effects of dosing volume and/or exposure times for CAIS agents. Two sets of experiments are conducted in Phase II. the first set, a small number of hairless guinea pigs are dermally dosed with the three agent solutions and with the neutralization solution using techniques for dosing chemical agents described in MREF SOP II-009 (enclosed). Using two animals at a time and a total of seven test sites per guinea pig, each animal is dosed percutaneously on both sides of the dorsal midline with low and high volumes of each CAIS agent mixture and with an optimal dosage of HD (approximately 1 μ L) to demonstrate the effect of a known vesicant. On the first dosing day, agent is allowed to remain in contact with the skin for approximately 2 hr (based upon results obtained in earlier MREF studies with hairless guinea pigs and HD), and the skin then decontaminated with an

approximately 0.5 percent sodium hypochlorite solution.* Approximately 24 hr following dosing, animals are sacrificed and skin samples from dosing sites are taken and fixed in formalin. Slides are prepared following tissue fixation, embedding, and sectioning, and evaluated for histopathology. Dosing volumes or times of exposure may be changed on subsequent days with additional animals to determine a volume and time of exposure for each CAIS agent mixture that results in consistent production of microvesication. Although the number of animals required to predict a dose volume and exposure time to consistently produce microvesication depends on the degree of microvesication observed in the first few animals, 4 to 12 animals are expected to be Once dosing volumes and times of exposure for each CAIS agent mixture are selected, the second set of experiments are conducted to verify consistent microvesication following administration of CAIS agent mixtures and to assess the extent of skin pathology following dosing of neutralization solution. For each CAIS agent mixture, dosing volume of neutralization solution is based upon the volume required to neutralize the volume of that agent mixture used to consistently create microvesication, but the maximum dosing volume at any site is limited to 100 μ L. Five quinea pigs are tested to verify microvesication in at least 80 percent of sites dosed with each of the CAIS agent mixtures and to ensure lack of microvesication at sites dosed with neutralization solution. If, for any of the CAIS components, either skin microvesication is observed on sites administered neutralization solution or the incidence of microblisters at sites dosed with CAIS agents is less than 80 percent, experimental procedures will be modified and the experiment repeated. Up to 10 animals will be used.

If problems are encountered in meeting the objectives described above, experiments will stop, the Contracting Officer's Representative (COR) and the ERDEC Task Area Manager (TAM) apprised of the situation, and an alternative approach agreed upon (with concurrence on significant changes by the ERDEC LAURC Chair). If feasibility and appropriate dosing parameters can be determined, the next phase will start.

Phase III: Evaluation of Efficacy of Neutralization Process

The objective of this final phase is to demonstrate that the neutralization process substantially reduces the

* In studies performed at the MREF, a 0.5 percent sodium hypochlorite solution has proven to be an effective decontaminant for HD-exposed animal skin without causing obvious irritation.

vesicating properties of CAIS agent mixtures. animal is percutaneously dosed with CAIS agent mixtures and with volumes of wastestreams using parameters established in Phase II. Dosing volumes are selected to contain equivalent agent quantities or maximum volumes of 100 μ L. Guinea pigs are sacrificed at approximately 24 hr after dosing and skin samples from dosing sites taken and prepared for histopathologic evaluation. It is anticipated that approximately 24 guinea pigs are required for this phase in order to demonstrate a statistically significant difference in the incidence of microvesication between sites treated with CAIS compounds and sites treated with neutralized CAIS compounds. If, after 12 animals have been dosed, a statistically significant (p < 0.01) difference is demonstrated, experimentation will cease at that time.

If problems are encountered in this phase, research will cease and the COR and TAM notified. Battelle and U.S. Army personnel will discuss problems and agree upon an alternative approach.

- B. Laboratory Animals Required and Justification:
 - Non-animal Alternatives Considered:

This task is necessary to determine if the vesicating properties of chemical agents have been virtually eliminated. This cannot be determined in other than a whole animal model.

2. Animal Model and Species Justification:

The hairless guinea pig is an appropriate and useful model to assess HD-induced vesication of skin (Marlow, et al., 1990; Mershon, et al., 1990). The hairless guinea pig bioassay model will be used to determine the extent of vesication before and after neutralization of agents.

- 3. Laboratory Animals:
 - a. Genus & Species: Cavia porcellus
 - b. Strain/Stock: Crl:IAF(HA)-hrBR
 - c. Source/Vendor: Charles River Lakeview
 (Newfield, NJ)
 - d. Age: Guinea pigs will be approximately 3 to 4 weeks of age upon receipt.
 - e. Weight: Guinea pigs will weigh approximately 200 to 350 g upon receipt.

- f. Sex: Male guinea pigs will be used in this study.
- g. Special Considerations: N/A
- 4. Total Number of Animals Required: 46 guinea pigs.
- 5. Refinement, Reduction, Replacement:
 - a. Refinement: Anesthetics will be used during exposure prior to decontamination.
 - b. Reduction: Experiments are conducted in a stage-wise fashion to limit the number of animals used to the minimum necessary to achieve statistically valid results. Procedures are stated for stopping experimentation if statistically significant results are obtained using fewer animals than expected to be necessary, or if problems are encountered. Results from previous studies at USAMRICD and at the MREF, and from previous phases of this study, will be used, as appropriate, to select doses and exposure times to limit the number of animals needed for this study to the minimum necessary to achieve statistically valid results.
 - c. Replacement: At the present time, vesication cannot be evaluated in other than a whole animal model.
- C. Technical Methods:
 - 1. Pain:
 - a. USDA (Form 18-3) Pain category:

Guinea pigs are anesthetized during the exposure period and it is believed that vesicating properties of the CSM will be virtually eliminated by neutralization and dilution. Positive control sites, i.e., sites dosed with CSM, have the potential to cause some pain, but animals are anesthetized during the exposure period. From past experience, guinea pigs do not appear to exhibit signs of pain following decontamination of vesicant agents. If signs of pain are exhibited following decontamination, buprenorphine at a dose of approximately 0.1-0.25 mg/kg sc can be given every 8-12 hr following consultation with a staff veterinarian or the study veterinarian.

- (1) No Pain
- (2) Alleviated Pain #46 100%

- (3) Unalleviated Pain or Distress
- b. Pain Alleviation:
 - (1) Anesthesia/Analgesia/Tranquilization:

Xylazine hydrochloride (approximately 6 mg/kg) and ketamine hydrochloride (approximately 35 mg/kg) will be given im to maintain anesthesia during exposure periods. Trained and experienced technicians will administer the anesthetics in the area of the hamstring muscles using a disposable tuberculin 1-mL syringe and a 23 to 25 ga needle.

- (2) Paralytics: N/A
- c. Alternatives to Painful Procedures:
 - (1) Source(s) Searched: TOXLINE, MEDLINE
 - (2) Date of Search: July 1995
 - (3) Key Words of Search: Mustard, Sulfur, Sulfur Mustard
 - (4) Results of Search: The hairless guinea pig model to measure microvesication as an indicator of dermal exposure to "blister agents" is the only well-accepted, published method to assess vesicating potential.
- d. Painful Procedure Justification: N/A
- 2. Prolonged Restraint: Restraint lasting more than approximately 2 hr is not anticipated, and guinea pigs will be anesthetized during this period.
- 3. Surgery: No surgery will be accomplished.
- 4. Animal Manipulations:

Guinea pigs selected for study are anesthetized with approximately 6 mg/kg xylazine hydrochloride and approximately 35 mg/kg ketamine hydrochloride or other veterinarian-approved anesthetic agent(s). Following anesthetization, animals are positioned in sternal recumbency on restraint boards. A maximum of eight dosing sites are demarcated using an indelible-ink pen. Within a chemical fume hood, guinea pigs are dosed percutaneously with CAIS agent(s), neutralization solution, wastestream samples, and/or control compounds depending upon the phase of study. Guinea pigs will

remain sedated/anesthetized during the time prior to decontamination with 0.5 percent sodium hypochlorite using additional doses of xylazine/ketamine (or other veterinarian-approved drug combination), as indicated. Following the decontamination of dosing sites, the animals are placed into individual polycarbonate cages within the hood. Use of Elizabethan collars may be necessary to prevent damage at dose sites. At approximately 24 hr following exposure, dose sites are evaluated for relative amounts of inflammation and pathology and approximation of size of any lesion. Guinea pigs are then euthanatized using deep inhalation anesthesia with halothane and death verified by opening the pleural cavity. Areas of skin at dosing sites are removed, placed in labelled cassettes, and put in a fixative solution. Tissue samples are processed and slides prepared by the Pathology Section of the Health Division or by MREF personnel and the slides are examined for histopathology and the presence or absence of microblisters by a qualified, experienced veterinary pathologist.

- a. Injections: Anesthetics only.
- b. Biosamples: No biological samples taken prior to necropsy.
- c. Animal Identification: Ear tags or tattoos will be used to maintain positive identification.
- d. Behavioral Studies: No behavioral studies will be done.
- e. Other Procedures: N/A
- 5. Adjuvants: N/A
- 6. Study Endpoint: The study will end approximately 24 hr following dermal exposure to test compounds when the animal is sacrificed and skin samples taken.
- 7. Euthanasia: Euthanasia will be accomplished by trained and experienced laboratory animal technicians under the supervision of a veterinarian. Guinea pigs are sacrificed using deep inhalation anesthesia with halothane followed by creation of pneumothorax.
- D. Veterinary Care:
 - Husbandry Considerations:
 - a. Study Room: Guinea pigs selected for study are anesthetized and positioned in sternal recumbency on restraint boards. Within a chemical fume hood,

guinea pigs are dosed percutaneously with CAIS agent(s), neutralization solution, wastestream samples, and/or control compounds depending upon the phase of study. Guinea pigs will remain sedated/anesthetized using additional doses of xylazine/ketamine (or other approved drug combination), as indicated, prior to decontamination with 0.5 percent sodium hypochlorite. Following the decontamination of dosing sites, the animals are placed into individual polycarbonate cages within the hood and held there overnight. Water and feed will be available ad libitum overnight.

- Special Husbandry Provisions: Hairless guinea pigs are held in isolation and observed for signs of clinical illness for at least 7 days prior to study initiation. Quarantine may be performed at Battelle's King Avenue Animal Resources Facility or at the MREF. All hairless guinea pigs are held at the MREF for at least 24 hr prior to study initiation. Hairless guinea pigs that are in apparent good physical condition after a minimum 7day quarantine period are selected for study. An ear tag or tattoo is applied for positive identification of each hairless guinea pig. Before being used in experiments, hairless guinea pigs are housed individually in stainless steel or polycarbonate cages equipped with a watering system. Fluorescent lighting with light and dark cycles of 12 hr each per day is provided. Room temperature of holding rooms is maintained at approximately 64-79 degrees F. At least 90 percent of the twice daily recordings will fall within the specified range. Relative humidity will be maintained at approximately 40-70 percent. At least 90 percent of the twice daily readings will fall within the specified range. Purina Certified Guinea Pig Chow® pellets are available at all times prior to study initiation. No contaminants which would interfere or affect the results of the study are known to be present in the feed. Analyses of the feed are maintained. Drinking water is supplied from the city of Columbus public water system at Battelle's Animal Resources Facility at King Avenue and from private wells when animals are housed at the MREF, and is available ad libitum prior to study. No contaminants which would affect the results of the study are known to be present in either water Water is analyzed annually for potability supply. and contaminants.
- 2. Attending Veterinary Care: Guinea pigs will be held for only approximately 24 hr following dermal exposures

before being sacrificed. Veterinarians are on staff and available for any emergencies which might arise.

- Enrichment Strategy: N/A
 - a. Dogs: N/A
 - b. Nonhuman Primates: N/A
- E. Data Analysis: For chemistry validation data generated in Phase I, tables of means and standard deviations of response of each control standard are prepared to present both the inter- and intra- variability of the analytical method. Calibration performance characteristics for each run, such as slope and standard error of the slope, R² (measure of fit about the regression line), method detection limits, and quantitation limits are presented in a table format.

For Phase II, inflammation and histopathology data for each dose volume and exposure time of each CAIS agent(s) are summarized and tabulated.

For Phase III data, statistical hypothesis tests are conducted at the 5 percent significance level to determine whether or not the neutralization process reduced the vesicating properties of agents contained in CAIS. For each CAIS ampule, incidence of microblisters at sites treated with CAIS agent(s) are compared to those of contralateral sites treated with the neutralized wastestream. Although incidence of microblisters is the primary endpoint for evaluating the efficacy of each neutralization process, analyses are also conducted on signs of inflammation, lesion area, and other histopathology data. To accommodate the intra-animal correlation of multiple measurements made on the same animal, McNemar's test or conditional logistic regression analyses may be used to analyze quantal data. Analysis of variance (ANOVA) models that include random effects for animal are fitted to continuous data. If data are not approximately normal, ANOVA may be conducted on transformed data, or nonparametric or categorical methods of analysis may be performed.

To minimize animal usage, Phase III experiments are performed using a two-stage, group sequential hypothesis test. The first stage consists of the experimental results for twelve animals. For each neutralization process, an interim analysis is performed using the data from these twelve animals. If microblister incidence at sites treated with wastestream is statistically less than that of sites treated with CAIS agent(s), then the evaluation is considered complete for that process;

sites previously used for that process may be employed for evaluating the remaining processes. Otherwise, up to twelve additional animals are tested, and the efficacy of the neutralization processes reassessed. The significance levels of the interim and final analyses are carefully controlled to maintain an overall type 1 error rate of 0.05. This may be accomplished by selecting significance levels of 0.01 and 0.05 for the interim and final analyses, respectively. The two-stage hypothesis test is conducted using the microblister data only.

- F. Investigator & Technician Qualifications/Training: The Study Director is an experienced research veterinarian and all animal technicians at the MREF are either AALAS certified as technicians or technologists or active in the AALAS training program. Records of their experience and training are available at the MREF.
- VI. Biohazard/Safety: Surety, security, and safety procedures for the use of chemical agents are thoroughly outlined in facility plans, in personnel requirements for qualification to work with chemical surety materiel (CSM), and in standard operating procedures for storage and use of CSM.

VIII. ASSURANCES:

- A. Animal Use: The animals authorized for use in this protocol will be used only in the activities and in the manner described herein unless an amendment is specifically approved by the IACUC.
- B. Duplication of Effort: I have made a reasonable, good faith effort to ensure that this protocol is not an unnecessary duplication of previous experiments.
- C. Statistical Assurance: I assure that I have consulted with an experienced, well qualified statistician in the design and strategy of this study, and the minimum number of animals needed for scientific validity will be used.
- D. Biohazard/Safety: I have taken safety into consideration in the design of this study and have made proper coordination in the preparation of this protocol.
- E. Training: I verify that the personnel performing the animal procedures/manipulations described in this protocol are technically competent and have been properly trained to ensure that no unnecessary pain or distress will be caused as a result of the procedures/manipulations.
- F. Responsibility: I acknowledge the inherent moral and administrative obligations associated with the performance of this animal use protocol, and I assure that all individuals associated with this project will demonstrate a concern for the health, comfort, welfare, and well-being of the research animals. Additionally, I pledge to conduct this study in the spirit of the fourth "R" which the DoD has embraced, namely, "Responsibility" for implementing animal use alternatives where feasible, and conducting humane and lawful research.

(Study Director)

G. Painful Procedures: I am conducting biomedical experiments which may potentially cause more than momentary or slight pain or distress to animals that will be relieved with the use of anesthetics. I have searched for alternatives to such procedures; however, I have determined that alternative procedures are not available to accomplish the objectives of the proposed experiment.

Can T Ma 11/29/95

(Study Director)

IX. Enclosure:

Battelle MREF Protocol II-009

X. References:

Marlow, D.D., Mershon, M.M., Mitcheltree, L.W., Petrali, J.P., Jaax, G.P., Sulfur Mustard-Induced Skin Injury in Hairless Guinea Pigs, J. Toxicol.-Cut. & Ocular Toxicol., 9(3), 179-192 (1990).

Mershon, M.M., Mitcheltree, L.W., Petrali, J.P., Braue, E.H., Wade, J.V., Hairless Guinea Pig Bioassay Model for Vesicant Vapor Exposures, Fund. and Appl. Toxicol., 15, 622-630 (1990).

MREF Protocol 109
Medical Research and
Evaluation Facility
January 24, 1996
Page 15
G1555-38A

Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components

Protocol Amendment No. 1

Change: Page 4, Section V. A. Experimental Design and General Procedures, Phase II:

Dosing Parameters for Agent(s) and Neutralization Solution: Change the wording
of the second sentence to read "In the first set, a small number of hairless guinea
pigs are dermally dosed with the three agents using the techniques for dosing
chemical agents described in MREF SOP II-009 (enclosed)." Change the wording
of line 11 from "volumes of each CAIS agent mixture" to "volumes of each CAIS
agent".

Page 5, Section V.A., Phase II: Change the wording on lines 9, 16, 19, 21, 23, and 27 from "agent mixture" to "agent" or "agent mixtures" to "agents".

Page 6, Section V.A., Phase III: Change the wording on lines 1 and 2 from "CAIS agent mixtures" to "CAIS agents". On lines 13 and 14, change "CAIS compounds" to "CAIS agents".

Page 11, Section V.E. Data Analysis, Change the second sentence of the third paragraph to read "Incidence of microblisters at sites treated with wastestreams is compared to that at sites treated with CAIS agent."

Reason for Change:

Instead of using actual ampules from CAIS kits, agent challenges will be prepared from HD, HN₁, and L stocks and will not be mixtures but individual agents. The neutralization solution will be dosed in the second set of experiments of Phase II, not the first as originally stated in the second sentence of the Phase II paragraph.

Change: Page 4, Section V.A. Experimental Design and General Procedures, Phase II:
Dosing Parameters for Agent(s) and Neutralization Solution: After the existing second sentence, as modified, add the following: "Because agents on a charcoal matrix require extraction prior to dosing, and because the ratio of neutralization solution to agent on charcoal is 70:1, the combined volume of agent and

MREF Protocol 109 Medical Research and Evaluation Facility January 24, 1996 Page 16 G1555-38A

neutralization solution for an agent dose that produces microblisters is greater than $100~\mu L$. The maximum dosing volume is $100~\mu L$. Therefore, it will not be possible to determine a volume of agent on a charcoal matrix that 1) produces microvesication and 2) can be neutralized and applied to a site. A sample of the wastestream from agents on charcoal will be tested for the ability to cause microvesication in Phase III, and will be chemically analyzed, but extracts of the agents on charcoal will not be tested in Phase II. Approximately 10 percent solutions of HD, HN₁, and L in chloroform, as well as neat HD, will be used to determine potential for creating microvesication. If, based on dosing volume limitations, one or two but not all three agents produce microvesication in the first set of Phase II experiments, the agent(s) producing microvesication in the lowest volume will be used to determine microblister formation following mixing of agent(s) with neutralization solution."

Reason for Change:

Because of limitations with dosing agents adsorbed on a charcoal matrix, it would be virtually impossible to administer a dose of agent which produces microblister formation combined with the neutralization solution in a total volume of less than $100~\mu L$. Therefore, agents in chloroform solutions, as well as neat HD, will be used to challenge animals. Those agents that demonstrate the formation of microblisters at the lowest dose volumes will be used to determine if the neutralization process substantially reduces the potential for creating microblisters.

Impact on Study:

These changes are not anticipated to have an impact on the study since they involve dosing only one agent rather than a mixture, and a 10 percent concentration is probably the maximum that could reasonably be expected for any CAIS agent other than neat HD.

MREF Protocol 109 Medical Research and Evaluation Facility January 24, 1996 Page 17 G1555-38A

Approved By:

Carl T. Olson, D.V.M., Ph.D.

Study Director

1/24/96

Date

Richard R. Stotts, D.V.M, Ph.D.

LTC, USA, VC

Contracting Officer's Representative

24 JAN96

Date

MREF Protocol 109
Medical Research and
Evaluation Facility
January 24, 1996
Page 18
G1555-38A

Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components

Protocol Amendment No. 2

Change: Page 1, Preface. Change to read:

"Study Performed by Battelle Memorial Institute's Medical Research and Evaluation Facility Building JM-3, West Jefferson, OH"

Reason for Change:

To clarify in the protocol the name and address of the testing facility where this study will be conducted, as required by 40 CFR Part 792.120 (3).

Change: Page 2, Sponsor. Add sponsor address so that this reads:

"Sponsor: Program Manager for Non-Stockpile Chemical Materiel (PMNSCM), USACMDA, Aberdeen Proving Ground, MD 21010-5425".

Reason for Change:

To add the sponsor's address to the protocol, as required by 40 CFR Part 792.120 (3).

Change: Page 2, I. Add, at the end of the paragraph, the following reference so that it reads:

"...Good Laboratory Practice (GLP) Standards (40 CFR Part 792)."

Reason for Change:

To add the specific reference for EPA GLP standards.

MREF Protocol 109 Medical Research and Evaluation Facility January 24, 1996 Page 19 G1555-38A

Change: Page 4, V.A., Phase I. Add at the end of the paragraph:

"The sponsor will provide wastestreams and HN. Composition of HD, L, and HN in CAIS also will be provided by the sponsor, and the components of the neutralization solution will be provided so that its preparation can be accomplished by MREF chemists."

Reason for Change:

To state compounds or documentation that will be supplied by the sponsor as required per 40 CFR Part 792.105 (a).

Change: Page 6, V.A., just prior to V.B., add:

"The first phase of this study will start after the receipt at the MREF of adequate quantities of HN and wastestream samples. Duration of the study is expected to be approximately 40 weeks to allow ample time for histopathologic evaluation of skin samples and analysis of results."

Reason for Change:

To add proposed experimental start and termination dates per 40 CFR Part 792.120 (4).

Change: Page 6, V.B.3.c., change to read:

"c. Source/Vendor: Charles River Laboratories."

Reason for Change:

The colonies of hairless guinea pigs needed for this study are raised at various locations and the source is not limited to the Lakeview colony.

Change: Page 14, IX. Change the word "protocol" to "SOP" so this reads:

"Battelle MREF SOP II-009."

MREF Protocol 109
Medical Research and
Evaluation Facility
January 24, 1996
Page 20
G1555-38A

Reason for Change:

This enclosure is an SOP, rather than a protocol.

Change: Page 14. Add section XI:

"XI. Records. Records to be maintained include, but are not limited to:

- A. CSM accountability log and inventory
- B. Chemical analyses and dose administration
- C. Animal data
- D. Clinical observations of lesions
- E. Histopathologic evaluations of lesions
- F. Decontamination, monitoring, and disposal records

Reason for Change:

This section was omitted from the original protocol and should be included per 40 CFR Part 792.120 (13).

Change: Page 14. Add section XII:

"XII. Reports.

- A. A draft final report will be prepared within 30 work days after completion of the exposures and analyses of the data. The draft final report includes:
- (1) Names of key study personnel
- (2) Experimental design
- (3) Test material description, analyses, preparation, and administration
- (4) Clinical observations
- (5) Histopathologic evaluations of skin samples
- (6) Statistical analyses of data
- (7) Discussions and conclusions.
- B. Following receipt of sponsor comments on the draft final report, a final report will be prepared within 30 work days.

MREF Protocol 109
Medical Research and
Evaluation Facility
January 24, 1996
Page 21
G1555-38A

C. Interim results will be reported to the COR verbally during the course of the study, as possible.

Reason for Change:

This section was omitted from the original protocol and should be included per 40 CFR 792.185.

Impact on Study:

These changes are not anticipated to have any impact on the study, but are meant to clarify portions of or add to the protocol and are primarily administrative in nature.

Approved By:

Carl T. Olson, D.V.M., Ph.D.

Study Director

2/6/90

Date

Richard R. Stotts, D.V.M., Ph.D.

LTC, USA, VC

Contracting Officer's Representative

Date

MREF Protocol 109 Medical Research and Evaluation Facility February 20, 1996 Page 22 G1555-38A

Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components

Protocol Amendment No. 3

Change: Page 8, V.C.1.b.(1), Anesthesia/Analgesia/Tranquilization. Change first sentence of this section to read:

"Xylazine hydrochloride (approximately 13 mg/kg) and ketamine hydrochloride (approximately 87 mg/kg) will be given im to maintain anesthesia during exposure periods."

Page 8, V.C.4. Animal Manipulations. Change first sentence to read:

"Guinea pigs selected for study are anesthetized with approximately 13 mg/kg xylazine hydrochloride and approximately 87 mg/kg ketamine hydrochloride or other veterinarian-approved anesthetic agent(s)."

Reason for Change:

After the first day of study, it was evident that a xylazine dose of approximately 6 mg/kg combined with a ketamine hydrochloride dose of approximately 35 mg/kg was not sufficient to induce an adequate stage of anesthesia for a suitable period of time. Optimal doses of xylazine and ketamine to induce general anesthesia in guinea pigs for 60 min has been reported to be 13 and 87 mg/kg, respectively.¹

Impact on Study:

This increase in dose of anesthetics should prevent the necessity of frequent booster doses of anesthetics and reduce any discomfort of the animals.

Wixson, S.K., Rabbits and Rodents: Anesthesia and Analgesia, in *Research Animal Anesthesia*, *Analgesia and Surgery*, Smith, A.C. and Swindle, M.M., Eds., Scientists Center for Animal Welfare, Greenbelt, MD, September 1994.

MREF Protocol 109 Medical Research and Evaluation Facility February 20, 1996 Page 23 G1555-38A

Approved By:

Carl T. Olson, D.V.M., Ph.D.

Study Director

2/20/96

Date

Richard R. Stotts, D.V.M., Ph.D.

LTC, USA, VC

Contracting Officer's Representative

20 Feb 96

Date

MREF Protocol 109 Medical Research and Evaluation Facility August 12, 1996 Page 24 G15555-38A

Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components

Protocol Amendment No. 4

Change: Page 6, V.A. Experimental Design and General Procedures: Phase III: Evaluation of Efficacy of Neutralization Process. Amend second sentence on this page, which reads "Dosing volumes are selected to contain equivalent agent quantities or maximum volumes of 100 µL." to read:

"Dosing volumes are selected to contain equivalent agent quantities, or are volumes of CAIS components and/or wastestreams selected by ERDEC, but do not exceed $100~\mu L$. If microvesication is observed following dosing with one or more wastestream, major fractions of wastestream(s) may be used to dose animals in efforts to determine whether parent compound or degradation products are causing lesions."

Reason for Change:

The "blue" wastestream, that CAIS component containing neat HD neutralized with DCDMH, has been found to cause microvesication in HGPs at dosing volumes that contain agent-equivalent amounts. Although not one of the recommendations for further studies suggested by the MREF, personnel of the PMNSCM office requested the dosing of additional animals with the "blue" wastestream and with the 10 percent HD in CHCl₃ solution at equal volumes. Additional studies may be performed with major fractions of the wastestream in an effort to determine the component(s) that are creating the microvesication.

Impact on Study:

This change should not affect the overall objectives of the study. Dosing of fractions of the wastestream should help the identification of components creating the microvesication.

MREF Protocol 109
Medical Research and
Evaluation Facility
August 12, 1996
Page 25
G15555-38A

Approved By:

Carl T. Olson, D.V.M., Ph.D.

Study Director

8/12/96

Date

Richard R. Stotts, D.V.M., Ph.D.

LTC, USA, VC

Contracting Officer's Representative

12 AUG96

Date

MREF Protocol 109 Medical Research and Evaluation Facility November 8, 1996 Page 26 G1555-38A

Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components

Protocol Amendment No.5

Change: Page 4, V.A. Phase I: Chemical Analyses and Validation of Methods

Change "Phase I: Chemical Analyses and Validation of Methods" to read "Phase I: Chemical Analyses and Evaluation of Methods."

Change sentence which reads "Each of the analytical methods, with any modifications, will be validated by evaluating the limit of detection, the limit of quantification, the linearity of response, and the precision, accuracy, and specificity." to "Each of the analytical methods, with any modifications, will be evaluated by determining the limit of detection, the limit of quantification, the linearity of response, and the precision, accuracy, and specificity."

Change the following sentence which reads "Analyses of the actual wastestreams for HD, HN and L concentrations will be accomplished following validation of the methods." to read "Analyses of the actual waste streams for HD, HN and L concentrations will be accomplished."

Page 11, V.E. Data Analysis

Change the first sentence from "For chemistry validation data generated in Phase I, tables of means and standard deviations of response of each control standard are prepared to present both the inter- and intra- variability of the analytical method." to read "For chemistry data generated in Phase I, tables of means and standard deviations of response of each control standard are prepared to present both the inter- and intra- variability of the analytical method".

Reason for Change:

Although the intent of Phase I of this study was to validate an analytical method for chemical vesicant agents in waste streams, no performance values were supplied to validate. Since this was the case, and because the provided method proved

MREF Protocol 109 Medical Research and Evaluation Facility G1555-38-A Page 27

inadequate for the intended purpose and a modified method was developed under a separate contract, a precision and accuracy analysis, rather than a validation, of the method was performed by the chemistry section of the MREF.

Impact on Study:

This change should not affect the overall objectives of the study. The analytical method originally provided proved to be inadequate, and modifications were made to the method. Precision and accuracy measurements of the modified procedure were accomplished to define the capabilities of the modified analytical technique.

Approved By:

Carl T. Olson, D.V.M., Ph.D.

Study Director

11/8/96

Date

Richard R. Stotts, D.V.M., Ph.D.

LTC, USA, VC

Contracting Officer's Representative

13 NOU90

Date

Battelle SOP MREF II-009-02 Page 1 of 9

STANDARD OPERATING PROCEDURE (SOP) FOR THE PERCUTANEOUS APPLICATION OF EITHER LIQUID OR VAPOR CHEMICAL SURETY MATERIEL OR OTHER IRRITANTS/VESICANTS ON SWINE AND GUINEA PIGS

Originated by:	Thomas A Frinder Thomas H. Snider, B.S.	Date <u>07-Feb-1</u> 99		
Approved by:	David L. Stitcher MREF Environment, Safety and Health Officer	Date 8Feh 96		
Approved by:	John B. Johnson D.V.M. Co-Principal Investigator and Manager Medical Research and Evaluation Facility	Date <u>PFeb</u> 96		
Reviewed and	Registered by QAU: Smulthalluta	Effective 2/96 Date 2/16/96		
Distribution Li	<u>st</u> :			
Quality Assurance Unit				

Battelle
Health Division
505 King Avenue
Columbus, Ohio 43201-2693

SOP Manual(s)

Battelle SOP MREF II-009-02 Page 2 of 9

I./II. Scope/Purpose

This standard operating procedure (SOP) describes all of the routine procedures for percutaneous application of sulfur mustard (HD), nitrogen mustard (HN), Lewisite (L), mustard Lewisite (HL) or other irritants/vesicants on swine or guinea pigs.

III. References

Battelle SOP MREF I-002, "Standard Operating Procedure (SOP) for the Storage, Dilution, and Transfer of GA, GB, GD, GF, TGD, VX, HD, HD/L, HN, and L When CSM Concentration/Quantity is Greater than Exempt Levels".

IV. Definitions

None

V. Procedures

A. Materials to be Used

CSM: HD, HL, HN, or L Other irritants/vesicants

B. Hazards Involved

Hazards are listed in Battelle SOP MREF I-002.

C. Handling of CSM

The handling of CSM is conducted in accordance with Battelle SOP MREF I-002. The procedures used within this SOP that are described in Battelle SOP MREF I-002 include: entry and hood set-up at MREF, obtaining, equilibration, transfer, dilution, transport, and securing vesicants.

D. Equipment

The following is a list of equipment which may be needed in addition to that listed in Battelle SOP MREF I-002: animal clippers and blades, gauze pads, skin decontaminant kits or decontamination solutions, scales, MINICAMS® or other head-space samplers, felt-tip pen, ruler, approved anesthesia solutions, slings, timer, approved euthanasia solutions, notebooks, blunt-tipped needles, microliter

Battelle SOP MREF II-009-02 Page 3 of 9

syringes, vapor cap assemblies, occluding material, o-rings, cyanoacrylate adhesive, Elizabethan collars, caging, underpads, tissue casettes, formalin solution, sharps container, warming pads/pumps, plastic bags, labels, scalpel handles and blades, volumetric pipettes, gas chromatography vials and caps, adsorbent material, large plastic jars with lids, dosing grid template, observation table, lab chair, calculator, disposable syringes, positive-displacement micropipettors and tips, trypan blue, tape, scissors, forceps, tongue depressors, bell jar, plastic-backed paper, brown kraft paper, holding boards, distilled water, cardboard pieces, surgical tape, and vet-wrap.

E. Preparation of Animals

Technicians working in the preparation room are, at a minimum, required to wear scrub suits, latex/nitrile gloves, and protective eyewear. If required by protocol, animals are anesthetized. Following initial anesthesia via injection or inhalation, booster anesthetic agents may be administered. A catheter may be inserted into a marginal ear vein and additional anesthetic provided intravenously. Booster doses of anesthetic may be administered to some species by intranasal drip or by inhalation. According to the procedures outlined in the specific protocol, the back of the animal may be cleaned prior to dosing. Dosing areas are marked as specified in the protocol. If required, Viton, Teflon, or rubber O-rings may be affixed to the back at dosing sites. If the protocol specifies removal of the animal from the hood on the day of dosing, double-sided tape is placed around the dosing area for affixing an air sampling container for performing proof-of-decontamination (POD). An animal may be secured to a restraint board (guinea pig) or in a sling (swine). Swine in slings will have their legs restrained to the posts of the stand. Animals are maintained in a fully anesthetized state while on restraint boards or in slings. Additional anesthetic agent will be readily available.

F. Preparation of Dosing Syringes

Syringes and blunt-tipped needles used for dosing will be chosen on the basis of the agent and the volume being applied. In general, the following guidelines are used.

Needles

Syringes

HD, HN - Stainless-steel
L - Platinum, iridium,
or gold-plated

< 0.1 μ L - Micrometer syringe device 0.1-1.0 μ L - syringe, e.g., Hamilton 7001

Battelle SOP MREF II-009-02 Page 4 of 9

HL - Platinum, iridium, or gold-plated
Others - As appropriate

1-10 μ L - syringe, e.g., Hamilton 701 10-100 μ L - syringe, e.g., Hamilton 1710 LT 10-500 μ L - syringe, e.g., Hamilton 1750 LT 100-1,000 μ L - syringe, e.g., Hamilton 1001 LT

The irritant/vesicant is drawn into the syringe following the procedure outlined in Battelle SOP MREF I-002. Once the syringe is properly filled, the syringe is placed on an adsorbent pad if more syringes are to be filled, or the content is expressed onto the designated dose site on the animal. The empty syringe is then placed on the adsorbent pad or reloaded for additional applications. After the appropriate number of syringes are filled, or the applications are made, a new lid is placed on the primary container.

G. Preparation of Dosing Micropipettors

A positive-displacement micropipettor may be used for the delivery of multiple aliquots. The doser should be aware that the potential exists for movement of the agent up the micropipettor neck and onto the surfaces of the mechanical parts inside the micropipettor. The doser should assume that such contamination has occurred, and following dosing, suspend the micropipettor on the rim of an Erlynmeyer flask containing distilled water to a level such that the plunger wire is submerged. After an approximately 24-hr period, the micropipettor may be dried and placed into a plastic bag and the bag sealed.

After each storage period and before use, the outer surface of the pipettor is examined for possible contamination. If there is any liquid on the exterior of the micropipettor, the pipettor is discarded by submersion in decontamination solution, and the doser removes and decontaminates his butyl gloves per SOP MREF I-002. If the micropipettor exterior appears uncontaminated, the micropipettor may be used. The doser examines the micropipettor to assure the proper volume setting.

The doser places the micropipettor on brown kraft paper or other disposable adsorbent surface, and picks up a plastic-backed wipe with one hand and a tip designed for that micropipettor in the other hand. The tip is surrounded by the wipe and held in one hand, the micropipettor is picked up in the other hand, and the tip put in place. The wipe is placed into decontamination solution. The tip is secured on the micropipettor. The tip diameter must be small enough to allow entry into the primary container of the CSM or irritant.

Battelle SOP MREF II-009-02 Page 5 of 9

The micropipettor is placed on brown kraft paper or other disposable adsorbent surface. The primary CSM or irritant container is uncapped per SOP MREF I-002. The micropipettor is picked up, and the tip inserted through the container opening until the tip point is just submerged. The micropipettor is never inserted to a position past the top of the disposable tip, which could cause contact of the micropipettor with the primary container's inside surface. The container is not appropriate for cosing with a micropipettor if the liquid level cannot be contacted without inserting the pipettor into the container.

The plunger is slowly allowed to retract, and agent or irritant is pulled into the tip. The viscosity of the material may require a pause to allow equilibration of pressure in the tip. The doser removes his thumb from the micropipettor plunger. The tip is withdrawn and wiped with a plastic-backed adsorbent paper which is then discarded into a beaker of decontamination solution. The doser moves the micropipettor to a position over the dosing site, and the plunger is carefully depressed to extrude the dose volume. If the used tip appears to be empty after dose delivery, it may be reused to perform multiple volume transfers. After the final use, decontamination solution is drawn from a beaker into the tip and dispensed into the waste beaker three times. The tip is removed and placed into a sharps container with decontamination solution.

H. Percutaneous Application of Liquid Irritant to Dose Sites

An individual wearing clean gloves will monitor the animal prior to dosing to ensure proper depth of anesthesia. If gas anesthesia is used, the anesthesia line will be secured prior to dosing. The volume of irritant applied to a particular dose site is protocol specific. Doses are applied to the center of each dose site using a syringe or pipettor with a blunt-tipped, positive-displacement needle. The tip of the needle may be touched to the application site to ensure transfer of material to the site from the syringe tip. Dermal applications requiring volumes larger than 5 µL may be made in incremental applications to the prescribed sites. Doses may also be applied as a streak. After dosing and any protocol-specified treatment and/or decontamination procedures, if applicable, are completed, experimental animals will be shifted within the hood system by the operations assistant. If possible, all animals will be positioned with their heads oriented toward the front of the hood. The animals will remain in this position until the end of the observation period. If the protocol requires holding guinea pigs within the hood system overnight, animals will be removed from their restraint and housed in appropriate caging with access to water. Feed may be made available if approved by protocol, the study director, or a staff veterinarian.

Battelle SOP MREF II-009-02 Page 6 of 9

I. Percutaneous Exposure by Vapor Cap Assembly

An assistant wearing clean gloves will monitor the animal to ensure proper depth of anesthesia. If gas anesthesia is used, the anesthesia line is secured prior to dosing. When the irritant/vesicant is administered as a vapor to an occluded dose site, it is dosed into a vapor cap assembly. The vapor cap assembly consists of a plastic cap with a smooth, flat rim of approximately 1 mm thickness, and a filter paper wafer placed inside the cap top. Carpet tape with adhesive on both sides is covered with release paper on both sides, appropriately sized openings for dosing made, and the tape cut into a size appropriate for mounting on the animal's back. Dose sites are marked on the animal's back as specified by protocol. If prescribed by protocol, a topical skin protectant (TSP) is applied to the dose site in an area slightly larger than the dosing perforation cut in the tape. The tape is applied to each dose site by removing the release paper from one side and firmly pressing that side onto the skin at the dose site.

Syringes or pipettors are selected based on the dose volume to be delivered and filled as described in Section F or G. The vapor cap assembly is placed upside down in a shallow well, approximately the same diameter as the cap and with a depth of approximately half the height of the cap, drilled into a holding block. At the time prescribed by protocol, the plastic film is removed from the top of the tape which is adhered to the skin at the dose site. The irritant is dispensed directly onto the paper wafer in the cap, taking care not to touch the needle or micropipettor tip to the rim of the cap. After the dose is delivered, the needle tip or micropipettor tip may be touched to the paper wafer, and the delivery device placed on a disposable adsorbent surface within the hood. The vapor cap assembly is then lifted out of the well with forceps, inverted, and placed onto a protocol-specified surface. After a designated time interval, each cap is transferred by sliding it onto a vapor cap transfer apparatus. The cap is then centered over the opening in the tape at a dose site and the cap top is lightly pressed with forceps to ensure a seal between the rim of the cap and the carpet tape. After receiving a particular treatment, animals are shifted within the hood system by the operations assistant. If possible, all animals are positioned so their heads are oriented toward the front of the hood. The animals remain in this position until the end of the observation period.

J. Decontamination of Irritant/Vesicant on Skin with a Test Decontaminant

Protocols may require the evaluation of test decontaminants on percutaneously dosed animals. The following describes the standard test decontamination

Battelle SOP MREF II-009-02 Page 7 of 9

procedure. Additional decontamination operations may be outlined within the protocol.

The dosing assistant or operations assistant applies a measured quantity of the test decontaminant to a swab (adsorbent padding, gauze, or fibrous pad secured to a tongue depressor(s)) or a decontamination mitt. At the scheduled time, the dosing assistant initiates decontamination by physically stroking or tapping, as specified by study protocol, the dosed area with a brisk action perpendicular to the animal's spine. For solid or powdered test decontaminants, a piece of cardboard may be held behind the dosing area to minimize sprread to other sites. After use, all swabs or mitts and other materials utilized for decontamination are deposited into a waste decontamination beaker. Quantity of test decontaminant to be applied, length of exposure before decontamination, and length of the decontamination process may be specified in the protocol.

K. Decontamination of Irritant/Vesicant following Vapor and/or Liquid Exposure

After a vapor exposure period, the vapor cap assembly is removed intact with the tape by gently pulling up on the folded tape tab with forceps. The vapor cap assembly and tape are placed into a bucket of the appropriate decontamination solution (e.g., 5 percent sodium hypochlorite (NaOCl)).

As specified by protocol and as directed by the sponsor, the dosed skin site may or may not be decontaminated following exposure. However, decontamination is necessary if the animal is to be removed from the hood.

In studies not involving TSPs, a decontamination assembly, consisting of either a 4 x 4 inch gauze pad or an adsorbent pad wrapped and taped around a wooden tongue depressor, is used to decontaminate the skin. A dry gauze pad is laid over the dose site for the protocol-specified length of time. An adsorbent pad assembly is moistened with a 0.5 percent NaOCl solution or other specified decontamination solution, and used to neutralize or remove any residual irritant from the dose site. The dose site is then rinsed using adsorbent assemblies moistened with distilled water. Once used, the assemblies are placed into decontamination solution.

If the protocol is for screening TSPs, a dry adsorbent may be used to wipe the TSP from the skin. The dose site is rinsed or treated as specified in the protocol. Any adsorbents used are placed into decontamination solution.

Battelle SOP MREF II-009-02 Page 8 of 9

L. Euthanasia, Decontamination and Removal of Animals

(1) Animals remaining in the hood overnight:

If specified by protocol, at the end of the observation period, the animals are anesthetized and an aqueous solution of trypan blue administered to aid in visualizing lesions. Upon completion of the experiment, all animals are euthanatized, either by administering an injection of a lethal dose of an approved euthanasia solution, or as specified by protocol. Tissues and/or skin samples may be taken if specified by protocol. The exposure site(s) and any possibly contaminated sites of all animals are treated with a forcepsheld gauze pad moistened with 5 percent NaOCl or other specified decontamination solution. Following the decontamination procedure, all gauze pads are discarded into a decontamination beaker.

Euthanatized and decontaminated animals are placed into a plastic bag. The plastic bag is brought to the hood face by a fully garbed technician. Outside the hood, a second technician awaits with a second or multiple layers of plastic bags, as specified. As the technician stationed outside the hood brings the bag(s) to the hood face, the technician working within the hood carefully places the bagged animals from inside the hood into the bag(s). The opening of each bag is twisted, folded, and taped. Labelled bags are brought to the POD area for analysis. All carcasses are to be incinerated after POD.

(2) Animals removed from the hood on the day of dosing:

A static air space over the dosing sites is monitored using air samplers. To achieve a static air space, a fitted flexible polyethylene container is secured to the animal's back over the dosing sites. The tightness of the seal is examined prior to initiation of sampling. (Note: The volume of the air space should be approximately 1 liter.) After the proper equilibration period, the cap from the sampling port is removed and a MINICAMS[®] sampler probe or other air sampling device is inserted into the static airspace. Animals cannot be removed from the hood until irritant/vesicant vapors are below established airborne exposure limits. When POD is obtained, clean gloves are donned, and the anesthetized animal is removed from the hood and placed into a holding cage. When the appropriate endpoint measurements have been made, the animal is euthanatized with an approved solution and, if applicable, skin samples are collected and placed

Battelle SOP MREF II-009-02 Page 9 of 9

into a formalin solution or other specified fixative. Animal remains are incinerated following the collection of skin samples.

M. Decontamination, Emergency Procedures, and First Aid Procedures

These procedures are described in Battelle SOP MREF I-002. All staff conducting these procedures are required to read, sign, and be familiar with Battelle SOP MREF I-002.

N. Quality Control

There are no procedure-specific quality control measures required for this SOP. When required, they are stated in the study protocol under which this SOP is performed.

DEVIATION REPORT

G155538A

Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components

Type of Deviation: SOP

During the compiling and quality control review of data generated during the conduct of the study, the Task Leader found that there were certain events that were performed not totally in accordance with an SOP. These events are listed below.

Date of Deviation: January 21, 1996.

Nature of Deviation: The relative humidity in Room 41 was recorded at 35% on this Sunday and there was no documentation of it being monitored, per MREF IV-001.

Cause of Deviation: Technician either did not realize the need to document or forgot to document additional readings which the SOP says will be taken when readings are outside the acceptable SOP range.

Impact on Study: None.

Corrective Action: Technicians working weekends will be reminded that "monitoring" out of range readings should be interpreted to mean that documentation of subsequent readings should be recorded during the time technicians are working in the facility.

Date of Deviation: March 5, 1996.

Nature of Deviation: The temperature and relative humidity conditions in Room 6, where the animals were prepared for dosing on this day, were inadvertently not recorded per MREF IV-001.

Cause of Deviation: Technician forgot to record these conditions.

Impact on Study: None.

Corrective Action: Technician was reminded to record these conditions per SOP. Also, when reviewing these records in December 1996 and January 1997, it should be noted that the SOP was revised in November 1996 to eliminate taking temperatures and humidities in non-animal rooms as this is not a necessary facility requirement. Duri Occause this technician no longer wasks at the MREF. Dam signing as her supervisor

Appendix A

Dates of Deviation: August 16, 1996 through August 26, 1996.

Nature of Deviation: The check for vermin in Room 17 was not documented per MREF VII-002.

Cause of Deviation: Technician forgot to document this check.

Impact on Study: None.

Corrective Action: Technician was reminded to document this check weekly. The

Date of Deviation: August 29, 1996.

Nature of Deviation: Room 6 was unavailable for animal preparation procedures so animals were prepared in another room which did not have a thermometer and hygrometer present per MREF IV-001.

Cause of Deviation: Staff member forgot to monitor these conditions.

Impact on Study: None.

Corrective Action: Staff member was reminded to record these conditions per SOP. Also, when reviewing these records in December 1996 and January 1997, it should be noted that the SOP was revised in November 1996 to eliminate taking temperatures and humidities in non-animal rooms as this is not a necessary facility requirement.

Prepared by:

A Marie Moore, Task Leader

Date

Date

2-13-97

Date

Approved by:

Carl T. Olson, Study Director

Date

Appendix A

Standard Operating Procedure (SOP) Deviation Report

Study Number: G1555-38A

Study Title: Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components

SOP Deviation to MREF III-002: SOP for the Measurement of Chemical Surety Materiel in Dilute Solutions of GA, GB, GD, TGD, GF, HN, HD/L, HD, L, and VX

Dates of Deviation: 2/20/96, 2/22/96, 2/27/96, 3/5/96.

Nature of Deviation: This SOP was revised in version -02, which became effective on March 6, 1996, to include the analysis of HN-1. Some HN-1 analyses were performed in February and early March on GLP study G1555-38A before version -02 was signed.

Cause of Deviation: Lag time in getting the new version approved and distributed.

Impact of Deviation on the Study: None. The methods used for HN-1 analysis are the same as those for HD, as stated in this SOP.

Corrective Action: None necessary.

Approved By: Carl T Date: 3/27/96

Approved By: Lichard Thill Date: 27 MAR 96
Study Sponsor

ANIMAL WEIGHT SHEET

G1555 -Study # <u>38</u>A

Project Number: 61555-38A

Species: H.G. Pig

Animal I.D. 30/ 302 30.3 30.5 306 307 309	Date: 2-2-96 Weight (kg) *** 304.6 3279 2901 3002 304.3 292.9
Animal I.D. 30/ 302 30.3 30.5 306	304.6 327.9 290.1 300.2
301 302 303 305 306	304.6 327.9 290.1 300.2
302 303 305 306	3279 2901 3002
303 305 306	2901 3002 3043
305 306	3002 3043
306	304,3
1 <i>307</i> 1	2929
309	305.8
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Tech. Init. Dina	

10 2-2-96

Note: Those aminds were assigned to a specific strake white several days after receipt. Therefore, initial or were not taken until 2-2= In order to help track the cut gain, they were again weighed on 2-796 and 2-16= Roberties

N/A = Not applicable

.00.09 500.09	
Balance Calibration: 100.09 500.09	Reviewed by: Robin Charles
Balance I.D.: / 53284	Date: 2-2-96
Weight I.D.: 211365	
Calibrated by:On_m	Dute: $\frac{\partial^2 - \partial^2 + \partial^2 \phi}{\partial x^2 - \partial x^2 + \partial x^2}$

Form No. MREF-WEIGHT.SHT-02.1 Appendix A

CD9 CD49 CHARLES RIVER LADORATORIES CD9 (PA)

CUSTOMER MASTER LABEL

8 IAF/HA-HO G. FIGS

250-300 GMS 23-29 DAYS

AREA: PO2

ACCDUNT #: 01825-01-041 PO #:115341

CONTROL # 1159454 SHIPMENT REF # 29612266

SHIPDATE: 02/26/1796

DELIVERY DATE:02/27/1996

TOTAL CRATES: 1

TRANSGEL PER CRATE: 1

008 01/31/1996

BATTELLE MEMORIAL INSTITUTE

INVESTIGATOR NAME: FRANCES REID SPECIAL REQUIREMENTS

CFW* MINIPIG-YU*

This sale is made on the terms on our current price list and no other.

Acceptance of this delivery constitutes your acceptance of those terms. MICROPIG* MINIPIG-HA*

8 hairless guirea pigs arrived at MREF at Approx. 12:30pm on 2-27-96. These were not originally ordered for Task 95-38, however, several days after their arrival, they were assigned for Task 95-38 use.

Animal Species: Heidess Guines fig Sex: M

Date: 2-27-96 Initials: Own.

.<u>.</u>..

Animal No.	Weight (g)	Male
499	261.3	
498	271.6	
497	250.0	
496	245.7	
494	275.8	
493	272.5	
492	275.0	
491	262.0	

These 8 hairless quien pigs arrived at the MREF on 2-27-96. On 3-1-96 it was decided that they would be assigned for use on Task 95-38.

Weights were taken on the day of receipt, however, a record of balance calibration checks, balance used, at was not documented at that time since those animals were originally ordered for a name of study. These animals were from the same source, all male, and received within the required weight range for Task 95-38 (G1555-38A).

Appendix A

102

3-1-96 Cal T Clow 10/30/96 Date:

: :

August 6, 1996

To:

G15538A Study File

From: Marie Moore, Task Leader

Subject: Under-Weight Hairless Guinea Pigs

For Protocol 109, during the first week of August, Charles River Laboratories was requested to send eight male Hairless Guinea Pigs weighing approximately 200-225 g on arrival so that the animals could be held for studies to be run later in the month and still not be excessively heavy. The protocol actually requires animals to weight approximately 200-350 g on arrival. On August 6, 1996, eight Hairless Guinea Pigs were received with three of the animals weighing under 200 g upon arrival. The study director is aware of this and does not anticipate any problem or adverse impact on the study.

Call T Dlsm 10/25/94

APPENDIX B

Analytical Methodology

Task 95-38 Analytical Method Verification G155538A

INTRODUCTION

This report summarizes the results of precision and accuracy testing of a gas chromatographic method with mass spectrometric detection (GC/MS) used for the determination of bis(2-chloroethyl)sulfide (HD), bis(2-chloroethyl)ethylamine (HN-1), and dichloro(2-chlorovinyl)arsine (L) in solutions neutralized by 1,3-dichloro-5,5-dimethylhydantoin (DCDMH) oxidation in organic solvent (Rapid Response System [RRS] neutralization). The analytical method was classified as "Interim" by the developer, but approved for use at Battelle's Medical Research and Evaluation Facility (MREF) by Edgewood Research, Development and Engineering Center (ERDEC) personnel. RRS neutralization samples were prepared by ERDEC personnel using the neutralization process for the method verification. These samples are referred to as waste streams or spent neutralization solutions in this report.

The RRS was designed by personnel from the office of the Program Manager for Non-Stockpile Chemical Materiel (PMNSCM) to neutralize vesicating compounds from field training kits (Chemical Agent Identification Sets) containing HD, HN-1 and L. These chemical warfare (CW) materiel are in organic solution (HD,HN,L), neat (HD), or adsorbed on granulated charcoal (HD,HN,L), and contained in sealed glass ampules. The analysis method designed requires a GC/MS system capable of splitless injection into a fused silica capillary column, analyte ionization by electron impact (EI), ion detection in the selected ion monitoring (SIM) mode, and a dedicated PC-based MS data processing capability.

METHODS

Method Summary

Method performance was determined through recovery from spent neutralization solution of agents added at known levels. Since the waste streams were expected to have residual oxidizing power, a direct spike into a sample waste stream, as received, would not be meaningful. Therefore, both the residual oxidant and the strong acidity (HCl) anticipated in the waste stream were quenched prior to the spike. Water soluble impurities were removed from the waste stream by partitioning to an aqueous buffer. In addition, L was derivatized prior to analysis by reaction with 1,3-propanedithiol (PDT) to form the arsenic disulfide derivative, 2-(2-chlorovinyl)-2-arsa-1,3-dithiocyclohexane [Cl-CH=CH-As(-SCH₂CH₂CH₂S-)], or L-PDT. The L-PDT derivative is referred to as L-Der in this report. An internal standard (IS), 1,2,4,5-tetrachlorobenzene, was used in agent quantitation and for GC retention time comparisons. Samples were analyzed without concentration of the extract, and quantitation was accomplished using calibration standards prepared in the extraction solvent mixture. Agent detection was assured for each waste stream sample through the recovery of an over-spike of the corresponding CW materiel within the sample matrix. Inadequate

recovery from spiked samples was presumed to be a result of sample matrix effects. These matrix effects were overcome by diluting and reanalyzing the sample.

The analytical method was developed by Dr. Samuel V. Lucas of Battelle's National Security Division, with the assistance of MREF chemists, under another task, and a copy of the method is attached to this appendixas Attachment A. The method was classified as "Interim" by Dr. Lucas, but MREF project staff were directed by ERDEC personnel to use the method.

Instrumentation

A Hewlett Packard Model 5970B Mass Selective Detector (MSD) was used for method verification and for subsequent waste stream analyses. The HP 5970B MSD is a stand-alone detector used in the MREF laboratory with an HP 5890A GC and the HP G1034C MS ChemStation for automating the analyses. The instrument conditions are presented in Attachment A.

Method Verification

There were three phases to the method verification.

Phase I. GC/MS Instrumental Performance with Calibration Standards

Calibration standards were prepared as mixtures containing approximately equal weights (μ g/mL) of HD, HN-1, and L-Der in the extraction solvent mixture at each analysis level. Triplicate preparations of each of five standards were prepared along with ten replicate dilutions of a 3 ppm standard. Initially, six standards were used but results indicated that the L-Der could not always be detected at or below 1 ppm so the 0.3 ppm standard was dropped from the data set. All three sets of standards and the ten separate dilutions of the 3 ppm standard were analyzed as a single sequence. The ten replicate injections from a single 3 ppm standard were included in the sequence to determine the instrument precision.

Phase II. Method Performance with Process Blanks

Process Blank is the term used in the method to indicate "control samples" taken through the entire sample work-up procedure. Seven separate spiked Process Blank extracts from each spike level - 2.5, 10, and 25 ppm (5 ppm was used for L instead of the 2.5 ppm due to the weak signal at 2.5 ppm) - were prepared and analyzed in a single sequence along with a set of five calibration standards and a Process Blank without agent.

Phase III. Method Performance with RRS Matrix (Waste Stream) Samples.

Three waste streams, identified below, were analyzed using the method. These samples were analyzed both as received (no agent spike) and in duplicate at each spike

level - 2.5, 10, and 25 ppm (5 ppm for L instead of 2.5 ppm). They were analyzed in a single sequence with a set of five calibration standards, and 0 and 25 ppm spiked Process Blank samples on three different days.

The three waste streams used in Phase III were identified as:

#4 Blue Process Fresh 5/96 lot # 96-0037-049 #5 Red Process Fresh 5/96 lot # 96-0037-047 #3 Charcoal Process 5/96 lot # 96-0037-014 L, -018 HN-1, -023 HD

These waste streams are referred to as Blue, Red and Charcoal in this report.

RESULTS

Each compound analyzed by MS has a characteristic fragmentation pattern, and this pattern along with the analyte's retention time provided a high degree of selectivity to the GC/MS technique. SIM data were acquired for the six ions specified in the method for each agent, and for the IS. Quantitation was achieved by using the target ion for each agent and the IS as specified in the method. The ratio of the peak area for the target ion of the analyte to the peak area of the target ion of the IS was used in the regression analysis. These quantities are referred to as target area and response ratio in the tables and figures of this report. The ChemStation analysis routine not only integrated the target ion peak area in the chromatogram, but also evaluated the relative response among three additional ions. In the ChemStation report, the retention time and target area were provided as well as the relative response of the three secondary, qualifying ions and an indication of whether the specified relative response criteria for the qualifying ions being investigated were met. On a few occasions when the analyte was absent (Process Blank or Waste Stream with no spike), the ChemStation integrator would report small target area values. In these cases a small shift in retention time was observed and the ChemStation report would indicate that the qualifications were not satisfied. Consequently, a value of zero was assigned as the target area by the analyst. The three dimensional data of time vs detector signal vs mass to charge ratio (m/e) add a significant degree of specificity to the method. The overall method specificity, however, was not evaluated in this study.

The SIM results for Phase I of the method verification are summarized in Table 1 of Attachment B (pages B-20 to B-22). The calibration curves for the agents are shown in Figures 1 to 3 on pages B-23 to B-25, and regression analysis results for the standards are presented in Table 2 on page B-26. Descriptive statistics for the three independent sets of calibration standards and the ten 3 ppm standards are summarized in Tables 3 and 4 on pages B-27 and B-28.

The SIM results for Phase II are summarized in Table 5 on pages B-29 to B-31. Regression analysis results for the standards analyzed with the Process Blank extractions are given in Table 6 on page B-32, and descriptive statistics for Phase II are summarized in Table 7 on page B-33

The SIM results for Phase III are summarized in Table 8 on pages B-34 to B-42. Regression analysis results for the standards analyzed with the waste stream extractions are given in Table 9 on page B-43, and descriptive statistics for Phase III are summarized in Table 10 on pages B-44 to B-49. A summary of the descriptive statistics for the waste stream samples over the three days of analyses are presented in Table 11 on pages B-50 to B-54. The percent recovery for each agent from the waste stream samples are presented graphically in Figures 4 to 6 on pages B-53 to B-55.

DISCUSSION

Phase I. GC/MS Instrumental Performance with Calibration Standards

Calibration fits were linear for HD and HN-1 and simple linear regression models were employed. A small but statistically significant quadratic effect was present for L-Der (Figures 1 to 3, and Tables 2 and 4). Therefore, all of the results reported for L-Der using calibration standards are based on a quadratic model. The regression models for all three sets of calibration standards were significant at the p<0.001 level, and r-squared values were all greater than 0.99. Statistical tests performed on the three independent sets of calibration standards demonstrated no significant differences between batches with respect to the regression fits of each analyte.

It was determined from preliminary data that reproducible results for the L-Der at 1 ppm were not practical. Therefore, the 3 ppm calibration standard was replicated to estimate the limit of detection and the limit of quantitation. The instrument limits of detection and quantitation were estimated with and without sample preparation. These values are reported in Table 3. Ten separate injections from a 3 ppm calibration standard along with a single injection from ten separate preparations of the 3 ppm calibration standard were used. The detection limit was computed as the concentration which corresponds to a signal that is three times the value of the standard deviation found for the lowest calibration standard. The quantitation limit is defined in a similar way using ten times the standard deviation. The detection limits found for the instrument were 0.4, 0.7, and 2 ppm for HN-1, HD, and L-Der, respectively. The quantitation limits for the instrument were approximately 1, 2, and 3 ppm for HN-1, HD, and L-Der, respectively (see Table 3).

Precision is evaluated as the relative standard deviation, which is reported as percent and is referred to as the coefficient of variation (C.V.) in the results tables. The precision of the instrument, as based on the analysis of calibration standards, is about \pm 5 percent for all of the analytes, and sample preparation adds about another \pm 1 percent (see C.V. in Table 3).

In Table 4, the relative standard deviation is given at each of the five concentration levels for the three independent sets of calibration standards. The value is 5 percent or less for all concentration levels for HD and HN-1. The relative standard deviation for L-Der increases with decreasing concentration so that at 1 ppm, the relative standard deviation is approximately 18 percent. Both the 1 and 2.5 ppm L-Der standards are less than the quantitation limit. Likewise, the error associated with the back-calculated concentration of the standards shown in

Table 4 are all well below 10 percent except for the two lower L-Der standards. The errors in the 0.8 and 8 ppm back-calculated concentrations for L-Der are both greater than 10 percent.

Phase II. Method Performance with Process Blanks

In Phase II, seven separate spiked Process Blank samples were extracted and analyzed at each spike level - 2.5, 10, and 25 ppm (due to sensitivity issues, a 5 ppm spike was used for L instead of 2.5 ppm). The relative standard deviations (C.V. in Table 7) were all less than 10 percent (range of 1.2 to 9.3 percent). At 25 ppm, all of the relative standard deviation values were less than 3 percent, indicating that the reproducibility of this multiple-step extraction process is good. The accuracy is expressed in percent recovery and is best for the 25 ppm: 105 percent for HD, 90 percent for HN-1, and 123 percent for L (Table 7). Since the percent recovery for 10 ppm HN-1 is much lower than that for 2.5 and 25 ppm HN-1, it was assumed that an error in preparing the solution used for this dilution had occurred. Each of the three spike levels were achieved by preparing three separate solutions and spiking each sample with 100μ L of the appropriate stock (both HD and HN-1 were mixed in a single solution at each level and L was prepared separately). Over the complete spiking range, the HN-1 concentration values (excluding the 10 ppm spiked sample discussed above) were underestimated by 10 to 15 percent. Except for the 25 ppm HD sample, which is 5 percent high, HD and L spiked concentration values were overestimated by 12 to 35 percent.

Phase III. Method Performance with RRS Matrix (Waste Stream) Samples

Low levels of HD or L were detected in all of the waste stream samples. The average concentration values over the three days were 12 ppm of HD in the Blue, 25 ppm of L in the Red, and 20 ppm of L in the Charcoal (Table 11).

The absolute recovery for the method was determined by spiking the waste stream samples. The three spike levels used in Phase II were used with each waste stream, and a 25 ppm spiked Process Blank sample was prepared. All of these samples were extracted with iso-octane for GC/MS analysis as outlined in the method. Calibration standards were analyzed sequentially with each set of extraction samples, and the regression analysis results for the calibration standards were used to determine the concentrations of HD, HN-1, and L in the initial 1-mL sample. Although L-Der is analyzed, the analysis reporting procedure calculates the concentration of L in the 1-mL waste stream or Process Blank sample.

The average percent recovery values over the three days ranged from 14 percent for the 25 ppm L spike in Blue to 143 percent for the 5 ppm L spike in Red. The 25 ppm spiked Process Blank sample was extracted and analyzed with the waste stream samples in Phase III to evaluate recovery efficiency. The recovery for these samples are given in Table 11 and summarized here.

Percent Recovery

	Process Blank	Blue	Red	Charcoal
HD	105	84	111	121
HN-1	8 6	62	94	8 6
L	138	14	104	105

The accuracy of the method was evaluated at the 25 ppm spike level by the difference between the percent recovery reported for the Process Blank and that of the various waste streams.

Method Accuracy

=	Blue	Red	Charcoal
HD	21 percent low	6 percent high	16 percent high
HN-1	24 percent low	8 percent high	0 percent error
L	124 percent low*	34 percent low	33 percent low

* Since the recovery of the 25 ppm L-spiked Process Blank was greater than 100 percent, the difference between the Process Blank and the waste stream sample is greater than 100 percent.

Only the 25 ppm results are discussed here. Results for the other two spike levels are provided in Table 11, and these values can be compared to the spiked Process Blank samples in Table 7.

In addition to percent recovery information outlined above, the mean and standard deviation of the back-calculated concentrations in ppm over the three days are given in Table 11. The standard deviation values for the spiked waste stream extraction samples were used to establish the method detection limits and the method quantitation limits as stated on page 4 for the instrument limits. The sample size for the spiked waste stream samples is 6 over the three days. The standard deviation values for all three spiking levels are approximately the same, so the average standard deviation over the 3 spiking levels was used. This combination makes the sample size greater than 10.

Detection and Quantitation Limits for the Method

	Bl	ue	Re	ed	Char	coal
	Method	Method	Method	Method	Method	Method
	Det. Limit	Quant. Limit	Det. Limit	Quant. Limit	Det. Limit	Quant. Limit
HD	3 ppm	10 ppm	2 ppm	5 ppm	4 ppm	14 ppm
HN-1	. 4 ppm	12 ppm	2 ppm	7 ppm	2 ppm	6 ppm
L	NR*	NR*	14 ppm	46 ppm	25 ppm	85 ppm

* NR - Not Reported. Since essentially no L-Der was recovered in the Blue waste stream, the estimated values would have little meaning.

Two sets of limits have been estimated because the calibration standards are prepared differently than the waste stream samples. The term instrument detection limit (or detection limit for the instrument) refers to the 99 percent confidence limit for detecting the analyte when it is prepared from a pure stock in the extraction solvent mixture like the calibration standards. The method detection limit refers to the 99 percent confidence limit for detecting the analyte when it is extracted from a waste stream matrix using the procedure in the method being verified. The method detection limits are always higher than the instrument detection limits because additional variables are introduced.

Conclusions

The extraction method evaluated offers a reproducible procedure (RSD < 12 percent at 25 ppm) for the analysis of HD and HN-1 in RRS waste streams. When the results are evaluated using the spiked Process Blank samples, spike recovery errors range between 0 and 24 percent at 25 ppm for HD and HN-1. The method detection limit and method quantitation limit vary with agent and waste stream, but they are roughly 5 and 15 ppm, respectively, for both HD and HN-1. Five and 15 ppm may be used as conservative estimates of the method detection limit and method quantitation limit, respectively, for all HD and HN-1 waste stream analyses. It is not possible to establish composite limits for L since the limits for the Charcoal waste stream are larger than those for the Red waste stream.

Calibration standards, with simple linear regressions, may be used for HD and HN-1, but a quadratic regression model is required for the analysis of L-Der. However, the reported concentration for L in the Red waste stream is near the method quantitation limit and the concentration for L in the Charcoal waste stream is near the method detection limit.

The RRS waste stream matrix interferes with the analysis process. This interference increases

the method detection limit and produces low recovery for L in spiked samples. The recovery of L is extremely low (124 percent lower than the 25 ppm Process Blank) in the Blue matrix (HN and L, however, are not present in the Blue neat HD ampules) and low in both the Red and Charcoal waste streams (34 and 33 percent at 25 ppm, respectively). A well trained operator is required for implementing this method, and judgments must be made about the character of the waste streams based upon the results. Accurate detection of L in a Blue waste stream is not possible, and underestimates of approximately 35 percent can be expected for both Red and Charcoal waste streams.

For agent recovery from Process Blank samples that are spiked at 25 ppm, there is an approximately 5 percent overestimated recovery for HD, 12 percent underestimated recovery for HN-1, and 30 percent overestimates of recovery for L (Tables 7 and 11). Except for HN-1, the recovery error increases at lower spike levels. Consequently, there are errors associated with the use of external calibration standards. The 25 ppm spike solutions were evaluated to assure that the recovery errors did not originate with the preparation of the spiked solutions. These solutions were diluted and analyzed without the extraction procedure. An independent set of calibration standards were prepared from the same original agent stock for the evaluation. The concentrations of the spike solutions were found to be within 4 percent of expected. During this analysis sequence, however, it was discovered that it was also necessary to correct for the concentration of the calibration standards prepared according to the method (see Attachment A, page B-18).

The stock solutions used throughout all phases of the Task 38 study, including those for dose confirmation and waste stream analyses, were prepared separately for each agent using chloroform as the solvent. There is no L-Der available for preparing calibration standards, so the L-Der was prepared from L stock as described in the method. The L-Der was washed with water to remove the HCl produced by the derivatization reaction and was then used as the stock for preparing calibration standards. With chloroform as solvent, it was not possible to wash the L-Der without loss. The first set of calibration standards was observed to have a large loss of L-Der in the final mixtures, so the first attempt to verify the method was not completed. After testing the L-Der preparation with just L, a second set of calibration standards was prepared using the procedure outlined in the method, and the verification process was repeated. In Phase II, during the second method verification process, it was again determined that the L-Der concentration was low. When a separate set of calibration standards was prepared (L-Der was separated from HD and HN-1, and the L-Der was not washed as specified in the method), the L-Der was determined to be 16.3 percent lower than expected. A 4.6 percent correction was also applied to the HD concentration during this process. Since there is no L-Der available for preparing calibration standards, these standards require careful evaluation.

One additional observation deserves brief attention. Low levels of L-Der, which could be

detected and quantified in calibration standards at the beginning of an analysis sequence, could not be detected in the same sample when injected later. Likewise, in Phases II and III of this study, the calibration standards were analyzed at the beginning and at the end of the sequence, and in all cases the response ratio for the standards at the later time were lower than the values recorded at the beginning of the sequence. This degradation in analysis system performance must be monitored closely by the analyst. For the results presented in this report, the difference in system performance adds a degree of bias within the sequence since the average response ratio values were used in the regression routine.

The analysis performance for L is not adequate for routine analyses. Quantitative measurements of L may be restricted to certain matrices and may require a time-based GC/MS injection specification.

ATTACHMENT A

Analysis Method

1. Title

Interim GC-MS method for the determination of bis(2-chloroethyl)sulfide (HD), bis(2-chloroethyl)ethylamine (HN-1), and dichloro(2-chlorovinyl)arsine (L) in neutralization mixtures based on 1,3-dichloro-5,5-dimethylhydantoin (DCDMH) oxidation in organic solvent (RRS neutralization system).

Note on the Classification of this Method as "Interim"

Two circumstances require this method to be designated as "Interim" at the present time: 1. The recovery of L spiked into RRS neutralization solutions used to treat high levels of HD ("Blue" RRS scenario) is generally very low or zero, apparantly due to a minor RRS system decon product of HD that can react with derivatized L. The interim solution for this problem is to dilute the matrix with RRS solvent and reanalyze spiked and non-spiked aliquots. 2. There is presently insufficient experience with method calibration in general as well as spike recovery over a wide range of RRS neutralization samples to predict that method instructions and performance specifications in these areas will not need modification.

While the "Blue" RRS process involves treatment of HD only (i.e., analysis capability for L in spent "Blue" neutralization solutions is irrelevant), it cannot be ruled out that field use of the RRS system might involve a high HD/low L scenario since the chemical identity of all field treatment items may not be known. Therefore, adequate method performance for all three CW materials (HN-1, HD, and L) in all RRS trial scenarios ("Blue", "Red", "Charcoal", and "Vermiculite") is considered essential for qualification of this method, and work to characterize and solve this L analysis difficulty is on-going.

In addition, a formal Level 1 P&A study has not been undertaken for this Interim Method. Therefore, all information relating to method detection limits, method quantification limits, analysis ranges, and other calibration and method performance and QC measures, while based on data available to date, must be regarded as likely to be revised as more experience and data from the application of the method is obtained.

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3. Kev Words

Rapid Response System, RRS, 1,3-Dichloro-5,5-dimenhylhydantoin, DCMH, neurtalization solution, Vesserting Agent, Sulfur Mustard, HD, Nitrogen Mustard, HN-1, Lewisite, L, Gas chromatography-Mass Spectrometry, GC-MS

4. Revision History

Current Revision: 10-Jun., '96 (DRAFT)

Previous Revisions: None

Original: 10-Jun., '96 (DRAFT)

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6. Purpose and Application

The Rapid Response System (RRS) has been designed by non-stockpile chemical materials (NSCM) to neutralize chemical agents sourced from field training kits containing HD, HN-1 and L. These CW materials are typically in organic solution or are neat, contained in sealed glass ampules. In some cases, broken ampoules will have contaminated surrounding vermiculite packing material which would also be subjected to the RRS neutralization system. One additional RRS matrix is granulated charcoal which is either known or suspected to have been exposed to one or more of these three CWs. The analysis method described here is designed to apply to all such applications of the RRS neutralization system to all of these matrices. The method requires a GC-MS system capable of splitless evaporator cavity injection into a fused silica capillary column, analyte ionization by electron impact (EI), ion detection in the selected ion monitoring (SIM) mode, and a dedicated PC-based MS data processing capability. A Hewlett Packard Model 5971A MSD with Chemstation II data system was used for the development of the method, but any GC-MS system with approximately equivalent capabilities should be sufficient.

7. Analysis Range

HD and HN-1 are thought to have an analysis (quantitation) range from ~1 to 100 µg per mL (ppm) of spent RRS neutralization solution with a detection limit between 0.1 and 0.5 ppm. The corresponding values for L are quantitation range of ~5 to 100 ppm and a detection limit between 0.5 and 2 ppm. A more accurate determination of these values will be made based on formal Level 1 P&A determinations planned for completion during Summer, 1996.

8. Summary of the Method

Method performance is documented for each sample through demonstration of the recovery from the spent neutralization solution of all three CW species spiked at an accurately known level, nominally 25 ppm. Because the anticipated spent neutralization solutions are expected to have residual oxidizing power, a direct spike into the sample, as received, is not meaningful. Therefore, both the residual oxidant and the strong acidity (HCl) anticipated in the spent neutralization solution is quenched prior to the spike. In addition, L must be derivatized prior to analysis by reaction with 1,3-propanedithiol (PDT) to form the arsenic disulfide derivative, 2-(2-chlorovinyl)-2-arsa-1,3-dithiacyclohexane [Cl-CH=CH-As(-SCH2CH2-CH₂S-)], i.e., L-PDT. An internal standard, IS, 1,2,4,5-tetrachlorabenzene, is used for CW quantitation as well as documentation of correct analyte GC retention. The sample is cleaned up by partitioning to aqueous buffer, and the extracted CW materials are analyzed without prior concentration of the extract. GC-MS analysis uses capillary gas chromatography, splitless sample injection, EI ionization, and SIM detection. Quantitation is against a calibration curve from GC-MS data for standards prepared in a simulated extract solvent. CW detection or non-detection is documented for each spent neutralization solution sample through the recovery of the corresponding CW material spiked into that sample. Inadequate recovery for these spiked samples requires dilution of the spent neutralization solution sample and re-analysis.

9. Materials and Equipment

a. Reagents and Chemicals

Reagent Water	KH₂PO₄	2,2,4-Trimethylpentane (Isooctane, i- C_s)
1,3-Propanedithiol (PDT)	K ₂ HPO₄	Thiolane (Tetrahydrothiophene)
1,2,4,5-Tetrachiorobenzene	tert-Butanol	Chloroform (HC stabilized)
Potassium Iodide (KI)	Acetic Acid	2,4-Dichloro-5,5-dimethylhydantoin (DCDMH)

In addition, SARM solutions of HN-1, HD and L are required to prepare calibration standards and solutions for spiking samples.

b. Equipment

GC-MS-Data System (HP 5971A MSD or other system with approximately equivalent or superior performance capability)

Disposable Glass Test Tubes, 16 mm x 125 mm (or functionally equivalent size), screw-cap with teflon cap liners

Syringes, 10 µL to 1,000 µL capacities

Solution Dispensers (three required), syringe pump type, up to 10-mL capacity (for dispensing 2.0 and 5.0 mL aliquots)

Pipettors and Tips (two required), 100 µL

Disposable Pipets, glass, 1.0 mL and 5.0 mL

Pipet Bulb

Disposable Transfer (Pasteur) Pipets, 5 % in. size

Pipet Pump for Pasteur Pipets (or Latex Bulbs)

GC Autosampler Vials and Caps (with teflon-faced liners)

Bench-top Centrifuge (with rotor accommodating test tubes, above)

Vortex Mixer

Glass Reagent Bottles, 1-liter, screw cap with plastic or teflon liner (for the conc. phosphate buffer) and 1- to 2-oz with teflon liner for the IS spiking solution)

Volumetric glassware for preparing solutions (100 mL graduated cylinder, 1,000 and 25 mL volumetric flasks)

c. Solutions

Concentrated Phosphate Buffer, 1.0 M in K₂HPO₄ and 0.6 M in KH₂PO₄ (RT storage)

Dilute Phosphate Buffer, prepared fresh each day by diluting Phosphate Buffer 1:20, v:v, with reagent water (do not store)

1,2,4,5-tetrachlorobenzene, 1.00 mg/mL in isooctane (i-C₃), this is the *IS* spiking solution (RT storage)

PDT in i-C₁, 1.0 vol.%

XDS (RDT&E) stock solutions of HD, HN-1 and L, unspecified solvent, (for preparing calibration standards and matrix spikes)

RRS Solvent: 48.5:48.5:3 (v:v:v), chloroform:tert-butanol:water (RT storage)

RRS Neutralization Solution: $\sim 0.55 \, \underline{M}$ DCDMH, 10.8 grams DCDMH dissolved to 100 mL final volume in RRS solvent ($\sim -10^{\circ}$ C or lower storage, 2-mo. storage life)

Water/Glacial Acetic Acid, 1:1, vol:vol

10. Sample Work-Up Procedure

(Note: "Sample" means either a spent neutralization solution aliquot or a process control (blank) aliquot; "vortex mix" means to briefly vortex mix and invert the tube at least twice—about a 5-sec procedure; samples are worked up in batches of 6 or fewer, depending on the centrifuge capacity with the work-up

procedure described below performed essentially simultaneously on all samples of the batch; see No. 11 for the definition of a "set" of samples.)

a) Approximate the relative oxidizing strength of the sample: Make a comparison standard by adding a carefully dispensed single drop (Pasteur (dispo.) Pipet) of full strength RRS neutralization solution into a test tube containing a measured 3.0 mL of 1:1, water:glacial acetic acid and 100 to 300 mg of KI (visually approximated; added and dissolved just prior to the assay). The orange color that develops is I₃ produced by the oxidizing capacity of the RRS neutralization system (DCDMH). Similarly assay the sample but continue to carefully add (counting) drops of sample until approximately the same orange color is achieved. However, do not add more than 20 drops. Record the approximate % original oxidizing power:

% [Ox] = (1/ # drops sample) x 100

If 20 drops still produce insufficient color density to match the reference color, approximate the fraction of reference color achieved to the nearest $1/5^{th}$ and adjust the %[Ox] value accordingly. For example, if the color density after 20 drops of the sample is 2/5 of the reference, % $[Ox] = 2/5 \times 5\% = 2\%$.

- b) Pipet 1.0 mL of sample into a test tube of the specified type; use of a glass pipet (not a pipettor) is recommended; if a pipettor is used, sufficient rinse steps must be performed until the pipettor will not drip sample within 5 seconds after filling (typically, at least 5 rinses). Chill the sample in an ice/water bath at least 3 min.
- c) Add 100µL thiolane (pipettor); vortex mix.
- d) Add 20 µL IS solution (25-µL syringe); vortex mix.
- e) Add 5 mL conc. phosphate buffer (dispenser pump); vortex mix.
- f) Add 100 µL PDT solution (pipettor); vortex mix.
- g) If the sample is to be spikes, add the spike aliquot at this point; add HD and HN-1 together or separately and L always separately and last; vortex mix after each spike addition.
- h) Add 2 mL i-C_s; vigorously shake all tubes in the batch simultaneously in a horizontal position for 60 timed seconds.
- i) Centrifuge just enough to cleanly separate the phases (typically, allowing the centrifuge to come to full speed and immediately shutting it off is sufficient); optionally, gravitational settling may be employed if both phases achieve clarity within -2 min or less which is typical for blank and diluted samples. If the organic layer is on the bottom, add 0.5 mL i-C₃ and repeat the shake-out and centrifugation (settling).
- j) Remove and discard the bottom (aqueous) layer using a dispo transfer pipet and the manipulator
- k) Add 5 mL dilute phosphate buffer (dispenser pump); shake (per step h) and centrifuge (per step i).
- l) Fill and seal a GC autosampler vial with the upper (i-C_s) phase; discard the remaining sample to the lab decon management system.

11. Grouping of Samples into Sets

An analysis "set" consists of samples and process blanks (including spikes) plus accompanying calibration standards that are subjected to GC-MS analysis in a single autosampler setup and which will have their data worked up together. No more than 6 RRS spent neutralization solution samples should be grouped into an analysis set. For each spent neutralization solution sample, an identical aliquot spiked at an accurately known level (nominally 25 ppm) with each CW agent is included. Also included are process blanks consisting of RRS Solvent in spiked and non-spiked forms and a duplicate spiked and non-spiked set for one of the spent neutralization solution samples of the set (regardless of the number of samples in the set). Thus, if only one RRS spent neutralization solution sample is to be analyzed, the set will consist of 6

sample workups: one each spiked and non-spiked process blanks, two non-spiked aliquots, and two spiked aliquots (i.e., duplicates) of the spent neutralization solution sample. Similarly, a maximum size set (6 RRS spent neutralization solution samples) would result in 16 sample workups.

12. GC-MS Analysis

No data are available on the storage stability of sample extracts. Therefore, the interim position is that sample extracts should be analyzed on the same day (or that evening) that they are generated. If this is not possible, store them at ~-10°C or lower no more than 5 days prior to analysis. Replace the injector liner at least after every 50 injections of spent neutralization solution sample extracts (not blanks or calibration standards) or sooner depending on the rate of sample residue and septum debris build-up in the liner. Use only wide-bore liners designed for splitless injection. Liners used with rapid injection autosamplers (HP 7673, for example) must be packed with deactivated glass wool.

GC Conditions	
Column	30 meter x 0.25 mm fused silica, 5% phenyl methylsilicone, 0.5 micron film (for example, Restek Rtx-5 #10238)
Carrier	Helium at 8 psi, pressure controlled
Purge (Splitter) Flow	30 to 50 mL/min
Injector Temp.	225°C
Transfer Line Temp.	275°C
Sample Injection	1.0 μL, splitless for 45 sec
Oven Program	65° (1 min hold); to 118° @ 15°/min (6 min hold); to 183° @ 12°/min (6 min hold); to 285° @ 25°/min (4 min hold). Total time is 30.03 min
MS Conditions	
Ionization	70 eV, Electron Impact
Detection Mode	Selected Ion Monitoring (SIM), see Table 1 for monitored ions
Ion Source/EM gain	HP autotune at "high sensitivity" or manufacturer specification equivalent
SIM Parameters	Monitor 6 ions for each species; for HP MSD, dwell time is 50 msec; for other instruments, select functionally similar SIM conditions.
Filament On Delay	6 min

Table 1. SIM Ions and Confirmation Ion Abundances

Species	Example ^(a) GC Retention, min	Quant. Ion / Conf. Ion (% of Quant. Ion)(b) / Aux. Ions
HN-1	9.8	120 / 1?2 (35%). 154 (4%), 134 (5%) / 92, 169
HD	10.6	158 / 109 (510%), 160 (68%), 111 (189%) / 123, 96
IS	14.9	216 / 214 (78%), 218 (48%), 181 (19%) / 220, 183
L-PDT	22.2	242 / 244 (35%), 181 (73%), 149 (117%) / 165, 107

⁽a) Actual GC retention will vary somewhat in the individual application; see discussion below.

⁽b) Values for "% of Quant. Ion" are modifiable based on the analyst's experience with calibration standards.

The GC oven program is designed to give optimal separation from potential interferences and adequate time to SIM changeover between HN-I and HD. The analyst may need to slightly vary the "hold" temperatures following the first and second temperature ramps in order to achieve elution of HD and L-PDT at the extreme end of their respective 6-min hold times. Alternatively, variation of the GC carrier supply pressure will have a similar effect. The requirement is that the peak top be located no earlier than one peak width nor later than two peak widths from the end of the respective 6-min hold period. The SIM ion changeover between HN-1 and HD should occur at a time nominally halfway between the two peak maxima.

13. Data Acceptance

GC-MS SIM Data—This method recognized that ion detection data acceptance always involves the analyst's judgment based on accumulating experience in the application of the method. However, this discretionary approach should operate within the following guidelines. In no cases are data to be accepted from spiked and non-spiked RRS spent neutralization solution samples without manual (i.e., direct visual) examination by the responsible analyst. This required data examination principally consists in verifying that peak areas appropriate to the level of CW material present are maximizing at the correct GC retention value and that they are present at approximately the correct ratios. The GC-MS data system can assist in this effort, but it cannot substitute for manual examination by the analyst. Often, this judgment must be made in conjunction with the analyst's recognition of overlapping or coeluting materials which interfere with the integrated peaks. In addition, the relative peak size criteria for confirmation and auxiliary ions (after accounting for apparent interferences) are expected to be less stringently met at lower levels of detection. For example, an auxiliary ion expected at 7% of the quantitation ion may be missing altogether for a bonafide detection which is supported by higher abundance confirmation or auxiliary ions. Final acceptance of data for the detection of HN-1. HD and L-PDT in the non-spiked aliquot for a given spent neutralization solution sample should always be justifiable by the chromatographic patterns observed for the corresponding spiked aliquot. Any analyst uncertainty in the qualification of data should be referred to the technical supervisor. Qualification of matrix sample data can be accomplished through hardcopy printouts of SIM chromatograms or by direct observation of them on the data system CRT. In the later case, data qualification which might be subject to dispute must de documented in hardcopy. This data acceptance process may result in reported detections which are below the P&A-generated method quantification limit; in these cases, the analyte is reported as "detected" but below the cited quantitation limit.

Spike Recoveries from Spent Neutralization Solution Samples—Spike recoveries from spent neutralization solution samples should be at least 80% for HN-1, 85% for HD, and at 70% for L and no greater than 110% for all three analytes. For HN-1 and HD, recoveries outside these ranges require a repeated sample workup and analysis (both spiked and non-spiked). In the event that the L spent neutralization solution sample spike recovery is less than 50% (as is expected for "Blue" scenario samples), dilute the sample 10x with RRS solvent and repeat the sample workup procedure and GC-MS analysis (both spiked and non-spiked). L analysis values for non-diluted samples with recoveries between 70% and 50% are adjusted for the actual L recovery found. For example, with the method quantification limit of L at 2 ppm, a sample with a non-detect on L and a 60% spiked L recovery would be reported as less than $2 \div 0.60 \equiv 3$ ppm; if the same sample (L spike recovery of 60%) resulted in L quantified at 35 ppm the reported value should be $35 \div 0.60 = 58$ ppm. For 10x-diluted samples, a similar adjustment is applied (in addition to the 10x dilution factor) regardless of the L recovery obtained. In all cases, samples which require dilution for L analysis must be reported as such.

14. Calibration

Calibration is based on a 5-point calibration curve using standards with analyte concentrations corresponding to those that would be obtained in the ~2.6 mL sample extract from 100% recovery of analytes present in the starting 1.0-mL sample at the following levels: 1, 3, 10, 30, and 100 ppm. Thus,

the actual concentrations of analytes in these calibration standards would not be 1, 3, 10, 30, and 100 ppm, but would be these values divided by 2.6. However, for convenience in preparing the calibration standards, a dilution factor of 2.5 is used. (Note, because quantification is based on an internal standard, this simplification is of absolutely no consequence for the correctness of the calibration obtained; also note that L is always detected as the derivative, L-PDT, but is quantified as the underivatized material). Thus the actual concentrations in the calibration solutions are 0.4, 1.2, 4, 12, and 40 µg/mL of CWAs and 8 µg/mL for the IS although the levels of identified to the data system are 1,3,10, 30, and 100 ppm of analyte and 20 ppm of IS in the starting neutralization solution sample. These standards are prepared in a solvent simulating the actual extract consisting of i-C₃:CHCl₃, 21:4, v.v. A stock solution of L-PDT is prepared from the L SARM in hydrocarbon solvent (hexane or i-C₃) by the addition of PDT at 5x the stoichiometric quantity and then washing the L-PDT once with unbuffered reagent water (to remove the HCl produced by the derivatization reaction). Once prepared, calibration solutions should be aliquoted to GC injection vials for storage at ~-10°C or lower until use. The use of 200 µL vial inserts to make multiple sets of calibration standards is recommended. At this writing, no data are available on the storage stability of calibration standards under these conditions.

Because it is not expected that this method will be in constant (daily) use, a calibration verification approach to data quantification is not used. Instead, a complete set of calibration standards are run with each set of samples (as defined in No. 10, above) as the first five analyses, and then the second-lowest standard is repeated at the end of the analyses. Data from the first five standards are used to generate a calibration curve to quantify the set (using the *IS* approach), and the final standard at the end serves as a calibration check standard (i.e., it is quantified along with the other samples).

The issue of acceptance of calibration data cannot be addressed at this stage because of insufficient experience in application of the method. However, it is anticipated that the curves for HN-1 and L-PDT will not be linear, and a quadratic or higher order fit will be necessary.

15. Quantitation

Quantitation of HN-1, HD, and L-PDT detected in samples employs the internal standard method using the peak areas for the quantitation ions listed in Table 1 and calibration curves generated from the calibration standards described in the preceding section. The internal standard method uses the ratio of the peak area of each analyte's quantitation ion to the corresponding value for the internal standard. It is assumed that the analyst responsible for applying this method is thoroughly familiar with the standard application of the internal standard method for quantifying chromatographic data. Analysts who are not experienced in this regard are not qualified to perform the role as the responsible analyst.

16. Reporting Results

Because analysis results are not meaningful without their accompanying quality control data, reports of analysis results should include the following and be reported as complete analysis sets (see No. 11 for the definition of an analysis "set"):

- a. The level found for each analyte in the non-spiked Process Blank and the spent neutralization solution samples included in the set. For the sample analyzed in duplicate, report the values obtained for each replicate as well as the average value.
- b. The relative residual oxidative power value for each spent neutralization solution sample.
- .. The percent recovery for the spiked Process Blank and the samples.
- d. The calibration equation and correlation coefficient for each analyte.
- e. The deviation (in %) of the end-of-set calibration check standard from its corresponding value with the other calibration standards.
- f. The approximate signal to noise level for the quantification ion in the lowest calibration standard.

g. A narrative description of any exception to the method taken with the set as well as any observations which would have interpretative value for the validity of the results obtained (i.e., unusual patterns of interference with detection ions or other observations in the GC-MS data or the behavior of samples through the sample extraction/cleanup procedure not usually encountered with similar samples).

17. Quality Control

The most important quality control issues are addressed by method provisions for the analysis of a spiked aliquot for every sample with provisions for minimum spike recoveries (No. 13) and for the generation of a complete calibration curve with each analysis set. Some additional quality control requirements are the following:

Record Keeping—The following records will be maintained in order to document the quality of the data produced

- a. A laboratory record book (LRB) will be used to record sample preparation activity including the preparation of standards and solutions and documentation of their storage conditions.
- b. A GC-MS log book will be maintained. All instrument use (including documentation of the analysis of sample sets, traceable to the sample preparation LRB) and maintenance and repair activity will be recorded in this log book. Typically documentation of analysis sample sets consists of a loose-leaf binder with printouts of the autosampler setup and file names and directory name and location of the data in the GC-MS data system.
- c. Hard copy print-outs of the mass spectrometer mass calibration report will be kept in a loose-leaf binder.

GC-MS System Maintenance—The GC-MS system manufacturer's recommendations on instrument maintenance are to be followed. In addition, criteria are given above for the routine changing of the GC injector liner. Decisions on the need for GC column replacement or removal of the inlet section are left to the judgment of the senior analyst responsible for the GC-MS data quality.

ATTACHMENT B

Results of Analytical Method Testing

TABLY, 1. SIM RESULTS FOR CALIBRATION STANDARDS

																	•	B-:	20																					-	
Response	ומרזם	0.0056	0.0189	0.0664	0.2159	0.6704		0.0060	0.0186	0.0639	0.2100	0.6802	0	0.00	0.0178	0.0635	0.1958	0.6985		0.0189	0.0186	0.0178	0.0177	0.0186	0.0366	0.0175	0.15	0.0159	0.0155		0.0189		0.0160	0.0166	55.00	6,10.0	0.016/	0.0160	0.0168	0.0168	0.0152
Tarret even		7,504	24,705	91,233	292,494	958,317		0/6//	24,533	85,472	282,477	929,955	7 946	696 66	53,333	81,481	266,266	960,011		24,705	24,533	23,353	23,310	25,219	22,327	23,992	20,114	20,088	19,843		24,705	21,291	19,884	20.590	20.894	20,651	106/07	20,043	100,02	876,02	18,913
Internal standard target area	' I	1,334,163	1,307,442	1,373,116	1,354,492	1,429,468	1,323.466	1 210 010	1,318,856	1,338,262	1,345,433	1,367,267	1,342,281	1.313.458	1 283 504	#6616971T	1,360,067	1,374,481		1,307,442	1,318,866	1,313,458	1,317,991	1,354,225	1,347,386	1,371,883	1,279,753	1,266,112	1,279,182		1,307,442	1,254,267	1,243,035	1,251,490	1,208,038	1,231,799	1.253.065	1.214.023	1,221,995	1 247 621	+ 1 - 1 1 1 - 1 +
Actual concentration (ppm)		96.0	2.88	9.64	28.8	96.4	96.0	α	0 64	40.7	28.8	96.4	96.0	2.88	9.64	• • • • •	20.0	96.4	ć	2.00	2.88	2.88	2.88	2.88	2.88	2.88	2.88	2.88	2.88		2,88	2.88	2.88	2.88	2,88	2.88	2.88	2.88	2,88	2.88	
Sample ID labe:		0/0296A5.D	070296A4.D	0.029683.0	070296A2.D	070296A1.D	070296B5.D	07029684.D	d E836C0C0	0.0000000	0/0296B2.D	070296B1.D	070296C5.D	070296c4.D	070296c3.D	n %296000	4.0000000000000000000000000000000000000	0,0295C1.D	G 6430000	4 14000000	0.029684.0	070296C4.D	0702964D.D	0702965D.D	0702966D.D	.0702967D.D	0702968D.D	0702969D,D	0702960D.D		070296A4.D	07029602.D	070296N3.D	07029 iJ4.p	07029605.D	07029606.D	07029607.D	07029638.D	07029509.D	07029610.D	
Calib. Std. Group		-					7						e	-					4	•										ı	ភ										
Compound	HD	2					HD						QH						QII											<u>.</u>											
	1	Αp	p∈	en	di	.x	В										1:	24																							

TABLE 1 (Continued)

			B-21		
Response ratio	0.0382 0.0997 0.3346 1.0648	0.0394 0.1018 0.3276 1.0436 3.3157	0.0986 0.3276 0.9747 3.3936	0.0997 0.1018 0.0986 0.0999 0.0940 0.0944 0.0921	0.0997 0.0921 0.0927 0.0946 0.0945 0.0911 0.0914
Target area	51,031 130,290 459,503 1,442,222 4,663,840	52,160 134,275 438,380 1,404,066 4,533,495	129,495 420,455 1,325,658 4,664,385	130, 290 134, 275 129, 495 131, 619 135, 133 126, 686 129, 448 117, 843 114, 380	130,290 115,506 115,206 118,330 114,183 115,391 114,141 110,930 116,171
Internal standard target area	1,334,163 1,307,442 1,373,116 1,354,492 1,429,468	1,323,466 1,318,866 1,338,262 1,345,433 1,367,267 1,342,281	1,313,458 1,283,594 1,360,067 1,374,481	1,307,442 1,318,866 1,313,458 1,317,991 1,354,225 1,347,386 1,371,883 1,279,753 1,279,753	1,307,442 1,254,267 1,243,035 1,251,490 1,208,038 1,231,799 1,253,065 1,214,023 1,221,995
Actual concentration (ppm)	1.01 3.03 10.1 30.3	1.01 3.03 10.1 30.3 101	3.03 10.1 30.3 101	3.03 3.03 3.03 3.03 3.03 3.03	3.03 3.03 3.03 3.03 5.03 5.03 5.03
Sample ID label	070296A5.D 070296A4.D 070296A3.D 070296A2.D 070296A1.D	07029685.D 07029684.D 07029683.D 07029682.D 07029681.D	070296c4.D 070296c3.D 070296c2.D 070296c1.D	070296A4.D 070296B4.D 070296C4.D 0702965D.D 0702966D.D 0702966D.D 0702969D.D 0702969D.D	070296A4.D 07029602.D 07029603.D 07029604.D 07029606.D 07029608.D 07029609.D 07029609.D
Calib. Std. Group	1	3 2		4	ហ
Compound	HN - 1	HN-1		- N - 1	HN-1
	Appendix	: B	125		

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			B	-22	
Response	0.0014 0.0084 0.0561 0.2718	0.0010 0.0084 0.0504 0.2576 1.1224	0.0010 0.0067 0.0430 0.2341	0.0084 0.0084 0.0067 0.0071 0.0075 0.0092 0.0056 0.0059	0.0084 0.0050 0.0059 0.0053 0.0055 0.0055
Target area	1,876 10,974 77,039 368,114 1,604,612	1,381 11,026 67,482 346,551 1,534,684	1,398 8,806 55,232 318,352 1,560,130	10,974 11,026 8,806 9,354 10,152 9,689 12,587 7,117 7,471	10,974 6,325 7,346 7,799 6,397 6,907 6,907 6,832 6,832
Internal standard target area	1,334,163 1,307,442 1,373,116 1,354,492 1,429,468	1,323,466 1,318,866 1,338,262 1,345,433	1,342,281 1,313,458 1,283,594 1,360,067 1,374,481	1,307,442 1,318,866 1,313,458 1,317,991 1,354,225 1,347,386 1,371,883 1,279,753 1,279,753	1,307,442 1,254,267 1,243,035 1,251,490 1,208,038 1,231,799 1,253,065 1,214,023 1,221,995
Actual concentration (ppm)	0.81 2.44 8.12 24.4 81.2	0.81 2.44 8.12 24.4 81.2	0.81 2.44 8.12 24.4 81.2	0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
Sample ID label	070296A5.D 070296A4.D 070296A3.D 070296A2.D	070296B5.D 070296B4.D 070296B3.D 070296B2.D 070296B1.D	070296c5.D 070296c4.D 070296c3.D 070296c2.D	070296A4.D 070296R4.D 070296C4.D 0702964D.D 0702966D.D 0702966D.D 0702968D.D 0702969D.D	070296A4.D 07029603.D 07029604.D 07029606.D 07029606.D 07029607.D 07029609.D 07029609.D
Calib. Std. Group	1	2	е	₹	r.
Compound	L-Der	L-Der	L-Der	L-Der	L-Der
	Appendix	с В	12	6	

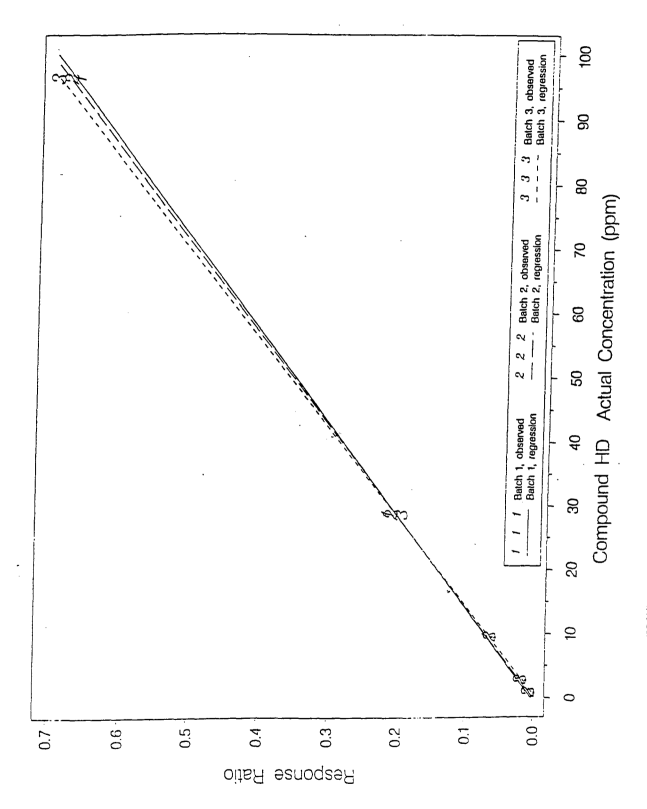
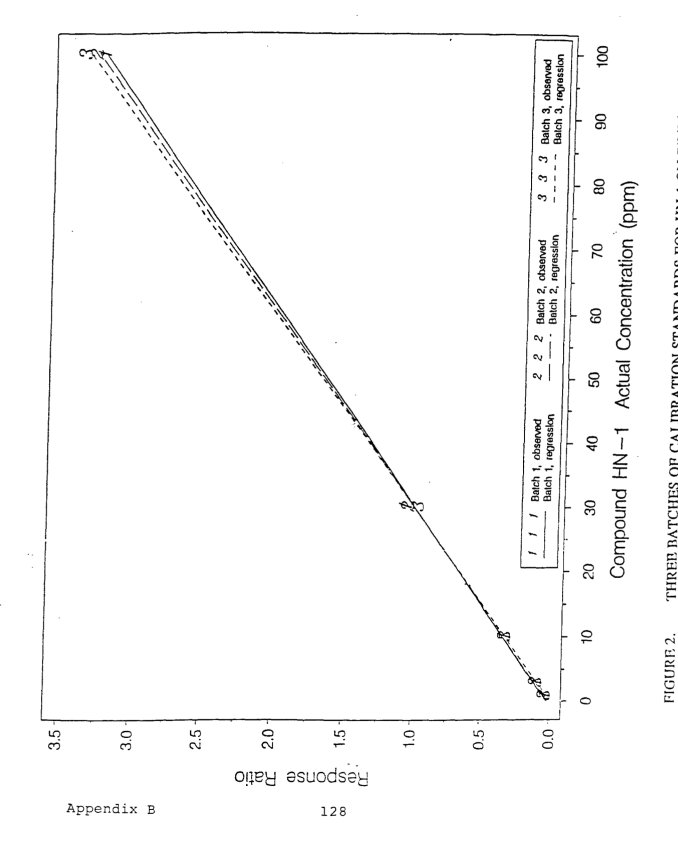


FIGURE 1. THREE BATCHES OF CALIBRATION STANDARDS FOR HD ON 7/2/96. OBSERVED RESPONSE RATIOS WITH FITTED LINEAR REGRESSIONS.



THREE BATCHES OF CALIBRATION STANDARDS FOR HN-1 ON 7/2/96. OBSERVED RESPONSE RATIOS WITH PITTED LINEAR REGRESSIONS.

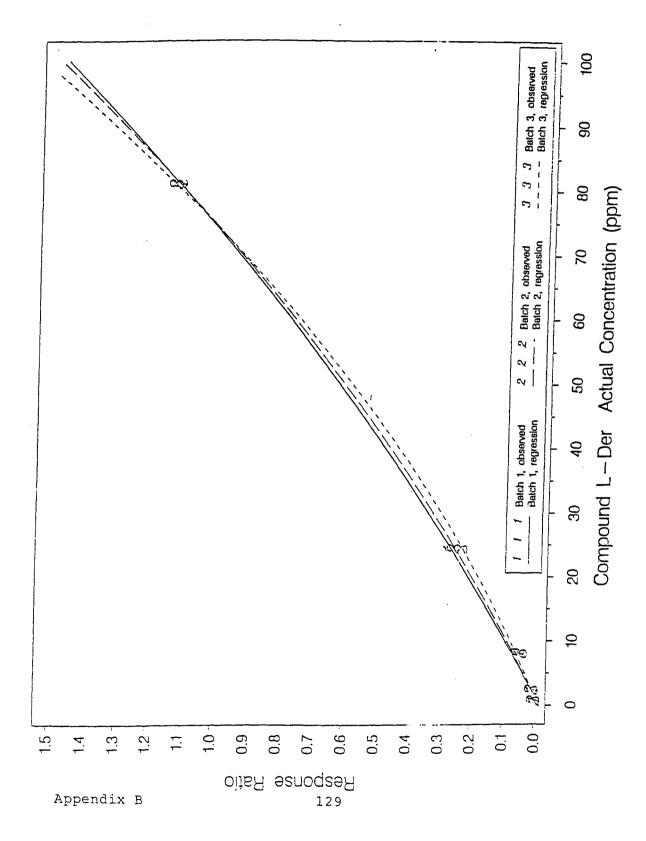


FIGURE 3. THREE BATCHES OF CALIBRATION STANDARDS FOR L.Der ON 7/2/96. OBSERVED RESPONSE RATIOS WITH FITTED QUADRATIC REGRESSIONS.

RESULTS FOR THREE INDEPENDENT SETS OF CALIBRATION STANDARDS IN THE 1-100 ppm RANGE TABLE 2.

	Compound	Batch	Intercept	Std.Err., Intercept	Slope coefficient	Std.Err., Slope	Slope2 coefficient	Std.Err., Slope2	Deviation from. Regression	
:	НD	1 2 3 All	0.002211 000604 005572 001322	0.004778 0.002595 0.003291 0.002390	0.00696751 0.00707831 0.00727766 0.00710782	0.000106 0.000057 0.000073 0.000053	1 1 1 1	, , , , , , , , , , , , , , , , , , , ,	0.008439 0.004583 0.005813 0.007311	
130	HN-1	1 2 3 All	0.021427 0.011112 012135 0.006801	0.024070 0.014161 0.012193 0.011246	0.03226638 0.03281458 0.03362845	0.000508 0.000299 0.000257 0.000237	1 1 1 1	1 1 1	0.042492 0.024999 0.021525 0.034387	D 26
	L-Der	1 2 3 A11	017109 016020 013961 015697	0.009859 0.009776 0.008822 0.005337	0:01053678 0.00964693 0.00812863 0.00943745	0.001029 0.001021 0.000921 0.000557	0.000043141 0.000053922 0.000074217 0.000057093	0.00001200 0.00001190 0.00001074	0.013552 0.013437 0.012126	

Regression lines or curves from the combined batches shown above as "All" are used to compute back-calculated calibration standard concentrations (ppm) shown in Tables 3 and 4.

Appendix

TABLE 3. DESCRIPTIVE STATISTICS OF RESPONSE RATIOS FOR EACH COMPOUND AND CALIBRATION SAMPLE TYPE GROUPS OF TEN CALIBRATION STANDARDS AT 3 ppm

Compound	Comparison standard (ppm)	Actual concentration (ppm)	z	Mean Response. Ratio	Back-calculated concentration (ppm) Hean S.D. C.V	d concentrati	on (ppm) c.v	Instrument Detection Limit	Instrument Quantitation Limit
HD HN-1 L-Der	е е е	2.88 3.03 2.44	10	0.0173 0.0963 0.0071	2.62 2.72 2.38	0.18 0.12 0.14	6.87 4.49 5.79	0.73 0.37 2.06	1.98 1.22 3.03 3.03
 		Sample	Name:	-Ten inject	- Sample Name=Ten injections from the same vial at 3 ppm	me vial at 3	wdd		
Compound	Comparison standard (ppm)	Actual concentration (ppm)	z	Mean Response Ratio	Back-calculated concentration (ppm) Mean S.D. C.V	ed concentrati	on (ppm)	Instrument Detection Limit	Instrument Quantitation Limit
HD HM-1	m m r	3.03	10	0.0167	2.54	0.14	5.46	0.60	1.57

TABLE 4. DESCRIPTIVE STATISTICS OF RESPONSE RATIOS FOR EACH AGENT AND CALIBRATION CONCENTRATION TABLE 1-100 ppm RANGE

Appendix

В						1	TOTAL WANGE		
		Comparison	Actual		Mean				
		standard	concentration		Response	Std.	Coef. of	Back-calculated	
	Compound	(mdd)	(wdd)	z	Ratio	Dev.	Variation	conc. (ppm)	
	HD	1	96.0	3	0.0059	0.0002	3.53	1.01	
		3	2.88	m	0.0184	0.0006	3.14	2.78	
		10	9.64	e	0.0646	0.0016	2.49	6.6	
1:		30	28.80	е	0.2072	0.0104	5.00	29.34	3
32		100	96.40	က	0.6830	0.0142	2.09	96.28	3-28
	HN-1	1	1.01	e	0.0378	0.0018	4.82	0 0	
		Э	3.03	Э	0.1000	0.0016	1.64		
		10	10.10	٣	0.3299	0.0041	1.24	£ 50.5	
		30	30.30	ස	1.0277	0.0471	4.58	20:0	
		100	101.00	3	3.3240	0.0658	1.98	100.82	
	L-Der	1	0.81	m	0.0012	0,0002	18.04	1 22	
		3	2.44	e	0.0078	0,0010	12.35	77.1	
		10	в.12	m	0.0499	0.0066	13.15	5. A	
		30	24.40	3	0.2545	0.0190	7.48	24 88	
		100	81.20	Э	1.1267	0.0073	0.64	0	

TABLE 5. RESULTS FOR THREE LEVELS OF SPIKED PROCESS BLANK SAMPLES

						B-2	9														
Response ratio	0.0293 0.0855 0.3155 1.0371 3.2203	0.000	0.0663 0.0638 0.0715	0.0611	0.0595 0.0596	0.2073	0.2039	0.2023	0.1828	0.1788 0.1989	0.7110	0.7175	0.6585	0.6968	0.7123	0.7104		0.0189	0.0650	0.05.0	3.0612
Target area	38,092 114,867 425,763 1,412,970 4,737,156	0	78,262 74,933 83,164	72,330	72,493 68,975	247,255	236,876	239,847 211,546	212,827	211,186 235,635	836,954	848,761	829,852	847,153	869,626	837,912		110107	191,988	1.310.296	4,650,198
Internal standard target area	1,301,818 1,343,718 1,349,641 1,362,438 1,471,008	1,276,293	1,180,442 1,175,005 1,163,004	1,182,918 1,160,199	1,218,560 1,157,195	1,193,019	1,161,619	1,185,399 1,154,956	1,164,549	1,180,942	1,177,204	1,182,894	1,260,301	1,215,836	1,220,911	1,179,453	1 350 604	100 000 T	1,379,819	1,365,829	1,519,061
Actual conc. of apike or std (ppm)	1.01 3.03 10.1 30.3	00.00	2.50	2.50	2.50	10.0	10.0	10.0	10.0	10.0	25.0	25.0	25.0	25.0	25.0	25.0	10.1		10.1	30.3	101
Sample Name	Calibration Std	Process Blank	Process Blank			Process Blank					Process Blank						Calibration Std				
Compound	11N-1	HN-1	HN-1			HN-1					HN-1						- 2	•			
	Appendix	В			1	L 3 3															

TABLE 5. (Continued)

								В	-30																		
Response ratio	0.0042 0.0138 0.0537 0.1891 0.6243	0.0000	0.0144	0.0148	0.0140	0.0136	0.0147	0.0120	0.0673	0.0668	0.0669	0.0666	0.0656	0.0652	0.0656	1131 0	0 1630	0.1611	0.1621	0.1628	0.1690	0.1637	8000	0.0028	0.0093	1/1/000	0.1/34
Target area	5,483 18,589 72,461 257,603 918,312	0	17,001	17,432	16,555	15,799	17,957	13,876	80,250	17,578	79,294	76,958	76,380	16,979	71,71	189,694	193.704	203,040	197,146	198,738	194,103	193,133	A . 7 9 A	12.160	601/21	239,519	941,300
Internal standard target area	1,301,818 1,343,718 1,349,641 1,362,438 1,471,008	1,276,293	1,180,442	1,175,005 1,163.004	1,182,918	1,160,199	1,218,560	1,157,195	1,193,019	1,161,619	1,185,399	1,154,956	1,164,549	1,180,942	1,184,756	1,177,204	1,182,894	1,260,301	1,215,836	1,220,911	1,148,545	1,179,453	1,350,594	1,308,467	1,372,819	1,365,829	1,519,061
Actual conc. of spike or std (ppm)	0.96 2.88 9.64 28.8	00.0	2.50	2.50	2.50	2.50	2.50	2.50	10.0	10.0	10.0	10.0	10.0	10.0	10.0	25.0	25.0	25.0	25.0	25.0	25.0	25.0	0.96	2.88	9.64	28.8	96.4
Sample Name	Calibration Std	Process Blank	Process Blank						Process Blank							Process Blank							Calibration Std				
Compound	HD	GI	HD						HD							OH.							НВ				
	Appendiz	х В						13	4																		

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	Response ratio	0.0026	0.0065	0.0463	0.2361	1.0404	0.000) 6250	0.550	0.0366	0.0322	0.0313	0.0312	0,0300		0.0756	0.0722	0.0712	0.0694	0.0670	0.0676	0.0670	0.2480	0.2463	0.2428	0.2413	0.2426	0.2442	0.2366	0.000	0.0017	0.0183	0.1331	0.8634
	Target area	3,324	8,788	62,554	321,725	1,530,502	o	44.701	42.142	42,512	38,035	36,278	38,013	34,695		90,209	83,885	84,415	80,191	77,973	79,839	79,365	291,988	291,390	306,053	293,402	296,243	280,528	279,005	0	2,171	25,139	181,746	1,311,522
Internal	brandard target area	1,301,818	1,343,718	1,349,641	1,362,438	1,471,008	1,276,293	1,180,442	1,175,005	1,163,004	1,182,918	1,160,199	1,218,560	1,157,195	; ; ;	1,193,019	1,161,619	1,185,399	1,154,956	1,164,549	1,180,942	1,184,756	1,177,204	1,182,894	1,260,301	1,215,836	1,220,911	1,148,545	1,179,453	1,350,594	1,308,467	1,372,819	1,365,829	1,519,061
Actual conc.	std (ppm)	0.81	2.44	8.12	24.4	81.2	00.0	5.00	5.00	5,00	5.00	5.00	5.00	5.00	ć	10.0	10.0	10.0	10.0	10.0	10.0	10.0	25.0	25.0	25.0	25.0	25.0	25.0	25.0	0.81	2.44	8.12	24.4	81.2
	Sample Name	Calibration Std					Process Blank	Process Blank							Jac la succession								Process Blank							Calibration Std				
	Compound	L-Der					L-Der	L-Der							7 - 7	יים מו							L-Der							L-Der				
		A	'bl	p e :	nd	lix	В						13	35																				

Appendix B

REGRESSION RESULTS FOR CALIBRATION STANDARDS ANALYZED WITH SPIKED PROCESS BLANK SAMPLES TABLE 6.

	Compound	Batch	Intercapt	Std.Err., Intercept	Slope coefficient	Std.Err. Slope	Slope2 coefficient	Std.Err., Slope2	Deviation from Regression	
1	91	1 2 Both	004052 009979 007016	0.002509 0.002981 0.002184	0.00652828 0.00651889 0.00652359	0.000055 0.000066 0.000048	1 1 1	1 1 1	0.004432 0.005264 0.005455	
.36	HN-1	1 2 Both	0.007645 009863 001109	0.022323 0.016669 0.021698	0.03196795 0.03052030 0.03124413	0.000471 0.000352 0.000458	1 1 1	1 1 1	0.039408 0.029427 0.054173	B-32
	L-Der	1 2 Both	014194 007591 010893	0.009218 0.005056 0.028695	0.00875368 0.00344704 0.00610036	0.000962 0.000528 0.002996	0.000052205 0.000089676 0.000070941	0.00001122 0.00000615 0.00003493	0.012671 0.006949 0.055781	

Regression lines or curves from the combined batches shown above as "Both" are used to compute back-calculated compound concentrations (ppm) shown in Table 7.

TABLE 7. DESCRIPTIVE STATISTICS FOR PROCESS BLANKS AND STANDARDS

В								
		Actual conc.		Mean				
		of apike or		Reaponse	std.	Coef. of	Back-calculated	Percent
Sample Name	Compound	etd (ppm)	z	Ratio	Dev.	Variation	conc. (ppm)	Recovery
Calibration Standard	GH	96.0	2	0.0035	0.0010	29.52	1.61	ı
		2.88	5	0.0116	0.0032	27.72	2,85	:
		9.64	7	0.0504	0.0047	9.30	8.80	1
		28.00	7	0.1822	0.0097	5.32	29.01	ı
		96.40	7	0.6220	0.0033	0.52	96.42	1
Calibration Standard	HN-1	1.01	2	0.0241	0.0073	30.45	0,81	ı
		3.03	7	0.0752	0.0145	19.30	2.44	ı
1		10.10	7	0.3003	0.0214	7.14	9,65	ŧ
37		30.30	7	0,9982	0.0550	5.51	31.98	B-
		101.00	2	3.1408	0.1125	3.58	100,56	33
	3		c	į	0	•	,	
Calibration Standard	r-ner	0.81	7	0,0013	0.0018	141.42	1.95	1
		2.44	7	0.0041	0.0035	84.19	2.39	t
		8.12		0.0323	0.0198	61.32	6.58	ı
		24.40	7	0.1846	0.0729	39.48	24.86	1
		81.20	7	0.9519	0.1252	13.15	81.18	,
Process Blank	HD	2.50	7	0.0139	0.0010	6.91	3.21	128.5
		10.00	7	0.0663	0.0008	1.21	11.24	112.4
		25.00	7	0.1634	0.0027	1.66	26.12	104.5
Process Blank	HN-1	2.50	7	0.0647	0.0051	7:90	2.11	. 84.3
		10.00	7	0,1939	0.0118	6.11	6.24	62.4
		25.00	7	0.7003	0.0202	2.88	22.45	89.8
Process Blank	L-Der	5.00	7	0.0336	0,0031	9.30	6.76	135.1
		10.00	7	0.0700	0.0032	4.60	11.68	116.8
		25.00	7	0.2431	0.0037	1.52	30.69	122,8

TABLE 8. RESULTS FOR THREE WASTE STREAM SAMPLES

Response ratio	ı	0.0331	0.0982	0.3693	1,1137	3,3488	0.000	0.0445	0.0420	0.1228	0.1215	0.4409	0.4171	0.000	.0801	-34		0.2130	0.7196	0.7539	0.000	0,0789	0.0935	0.2055	0.1914	0.6707	0.7073	0.000	0.6694		6610.0	0.0664	0.2812	0.9653	.0005
Re Target area r		40,734					0			165,463			547,966	0		,950		234,215			0	87,397	100,531	222,253			756,576	0	705,033					e	3,653,239 3.
standard target area		1,229,757	1,253,857	1,265,599	1,269,552	1,363,286	1,419,824	1,316,665	1,315,639	1,347,568	1,288,883	1,280,120	1,313,609	1,206,279	1,099,030	1,077,885	1,142,997	1,099,497	,101,	1,054,513	1,175,556	1,107,506	1,075,616	1,081,556	1,107,330	1,100,883	1,069,669	1,107,402	1,053,244	1.146.750	1 160 400	1,150,405	1,158,131	1,155,481	1,217,546
of apike or atd (ppm)		1.01	3,03	10.1	30.3	101	00.00	2.50	2.50	10.0	10.0	25.0	25,0	0.00	2.50	2.50	10.0	10.0	D.	25.0	00.00	2.50	2.50	10.0	10.0	25.0	25.0	0	25.0	1.01	3 03	7	10.1	30.3	101
Waste Stream							Blue							Red							Charcoal							Process Blank							
Sample Name		Callbration Std					Extraction Sample							Extraction Sample							Extraction Sample							Extraction Sample		Calibration Stc					
Injectinn Date		07/08/95					86/80/10							96/80/10							96/80/10							96/80/10		96/80/10					
Compound		T-NE A	pp	oe i	nđ	ix	B 188-1							HN-1	,	13	8				IIN-1							HN-1		HN-1					

TABLE 8. (Continued)

07/10/96 Calibration Std 1.01 1,311,822 36,106 0.07075 10.01 1,541,779 1.12,879 0.01046 10.01 1,628,767 1.12,899 0.01046 10.01 1,628,767 5,134,994 1.2755 10.00 1,628,743 6.156,744 0.0000 2.50 1,556,444 79,736 0.0508 2.50 1,566,744 79,736 0.0508 2.50 1,566,744 79,736 0.0508 2.50 1,566,744 79,736 0.0508 2.50 1,566,744 79,736 0.0508 2.50 1,566,744 79,746 0.0000 2.50 1,566,744 79,746 0.0209 2.50 1,566,744 79,749 0.0209 2.50 1,566,744 79,749 0.0209 2.50 1,393,992 112,499 0.0499 2.50 1,317,402 1.22,499 0.0209 2.50 1,317,402 1.05,797 0.0879 2.50 1,317,402 1.05,797 0.0879 2.50 1,317,109 0.0000 2.50 1,317,109 0.0000 2.50 1,317,109 0.0000 2.50 1,317,109 0.0000 2.50 1,317,109 0.0000 2.50 1,317,109 0.0000 2.50 1,317,109 0.0000 2.50 1,317,109 0.0000 2.50 1,317,109 0.0000 2.50 0.0101 2.50 1,317,109 0.0000 2.50 0.0101 2.50 1,317,109 0.0000 2.50 0.0101 2.50 1,317,109 0.0000 2.50 0.0101 2.50 1,317,109 0.0000 2.50 0.0101 2.50 1,317,109 0.0000 2.50 0.0101 2.50	Compound	Injection Date	Sample Name	Waste Stream	Actual conc. of spike or std (ppm)	internai standard target area	Target area	Response ratio	
10.3 1,502,948 1,562,757 1,0358 1,035	1-1	07/10/96			1.01 3.03 10.1	1,311,822 1,452,174 1,541,279	36,106 122,879 478,512	0.0275 0.0846 0.3105	
07/10/96 Extraction Sample Blue 0.000 1,650,237 0.0000 0.0538					30,3	1,502,948 1,628,767	1,562,767 5,334,994	1.0398 3.2755	
2.50 1,576,644 79,736 0.0506 10.0 1,444,227 228,680 0.1441 10.0 1,444,227 228,680 0.1441 25.0 1,566,114 786,266 0.5020 25.0 1,566,114 786,266 0.5020 25.0 1,566,114 786,266 0.5020 25.0 1,393,392 122,449 0.0867 25.0 1,393,392 122,449 0.0867 25.0 1,314,419 124,377 0.0867 25.0 1,312,407 127,737 0.0867 25.0 1,312,407 127,737 0.0867 25.0 1,317,64 102,155 0.0819 25.0 1,317,64 102,155 0.0819 25.0 1,317,64 102,155 0.0819 25.0 1,317,64 102,155 0.0819 25.0 1,317,64 102,155 0.0819 25.0 1,247,64 102,155 0.0819 26.0 1,247,64 102,155 0.0819 27.0 1,247,64 102,155 0.0819 28.0 1,247,64 102,155 0.0819 28.0 1,247,64 102,155 0.0819 28.0 1,247,64 102,155 0.0819 28.0 1,247,64 102,155 0.0819 28.0 1,247,64 102,155 0.0819 28.0 1	HN-1	07/10/96	Extraction Sample		0.00	1,650,237	0 83,560	0.0000	
10.0 1,544,227 228,680 0.1444 25.0 1,566,114 786,262 0.5020 25.0 1,566,114 786,262 0.5020 25.0 1,566,114 786,262 0.5020 25.0 1,566,114 786,262 0.5020 25.0 1,393,392 122,497 0.0607 25.0 1,394,419 124,377 0.0607 25.0 1,394,428 124,377 0.0607 25.0 1,314,428 124,377 0.0607 25.0 1,314,428 124,377 0.0607 25.0 1,314,428 124,377 0.0607 25.0 1,314,428 124,377 0.0607 25.0 1,314,409 124,377 0.0607 25.0 1,314,409 124,377 0.0607 25.0 1,314,409 124,377 0.0607 25.0 1,314,409 124,377 0.0607 25.0 1,314,409 124,377 0.0607 25.0 1,314,409 124,377 0.0607 25.0 1,240,262 1,039,955 0.0693 25.0 1,240,262 290,664 0.2344 25.0 1,240,262 290,664 0.2344 25.0 1,240,262 290,664 0.2344 25.0 1,211,240,262 290,664 0.2344 25.0 1,211,296 955,995 0.6971 25.0 1,212,21 8658,465 0.0191 25.0 1,212,221 8658,465 0.0191 26.0 1,216,775 167,024 1.0323 26.0 1,216,775 1470,024 1.0323					2.50	1,576,644	79,736	0.0506	
25.0 1,566,114 786,266 0.5020 25.0 1,566,627 817,632 0.5253 07/10/96 Extraction Sample Red 0.00 1,532,021 0.00879 2.50 1,393,392 122,449 0.0879 2.50 1,314,428 315,281 0.2439 10.0 1,314,428 315,281 0.2439 10.0 1,313,184 1,061,874 0.0805 2.50 1,313,184 1,039,955 0.0819 2.50 1,313,184 1,061,874 0.0805 2.50 1,313,184 1,039,955 0.0819 2.50 1,313,159 106,876 0.0868 2.50 1,213,159 106,876 0.0819 10.0 1,215,439 281,071 0.2131 25.0 1,179,525 816,801 0.7094 25.0 1,317,298 955,995 0.6971 07/10/96 Extraction Sample Process Blank 0.00 1,215,439 0.0000 25.0 1,317,298 0.0001 25.0 1,317,298 0.0001 25.0 1,317,298 0.0001 25.0 1,317,298 0.0011 25.0 1,317,298 0.0011 25.0 1,317,291 0.0001 25.0 1,317,291 0.0001 25.0 1,317,291 0.0001 25.0 1,317,291 0.0001 25.0 1,317,291 0.0001 25.0 1,317,291 0.0001 25.0 1,317,291 0.0001					10.0	1,587,228	229,234 228,680	0.1444	
25.0 1,556,627 817,632 0.5253 07/10/96 Extraction Sample Red 0.00 1,532,021 0 0.0000 2.50 1,334,428 124,377 0.0867 10.0 1,314,428 135,381 0.2439 10.0 1,312,407 337,377 0.2439 10.0 1,312,407 337,377 0.2439 25.0 1,312,407 337,377 0.2439 25.0 1,312,407 337,377 0.2439 25.0 1,313,100 0 0.0000 2.50 1,217,644 1,061,874 0.00188 2.50 1,217,644 102,155 0.0819 10.0 1,217,443 139 281,071 0.7094 25.0 1,217,439 255 0.0819 25.0 1,215,439 0.0000 25.0 1,217,439 0.0000 25.0 1,217,221 888,465 0.0313 25.0 1,226,030 0 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,634 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000 25.0 1,218,439 0.0000					25.0	1,566,114	786,266	0.5020	
07/10/96 Extraction Sample Red 0.00 1,532,021 0 0.0000 2.50 1,393,392 122,449 0.0879 2.50 1,394,418 124,377 0.0867 10.0 1,314,418 135,281 0.2439 10.0 1,312,407 327,737 0.2439 25.0 1,312,407 327,737 0.2497 25.0 1,312,407 327,737 0.2497 25.0 1,311,100 0 0.0000 0.00000 1,311,100 0 0.00000000000000000000000000000					25.0	1,556,627	817,632	0.5253	
2.50 1,393,392 122,449 0.0879 2.50 1,434,419 122,449 0.0867 10.0 1,744,419 135,281 0.2439 10.0 1,312,407 327,737 0.2439 10.0 1,312,407 327,737 0.2439 2.5.0 1,323,184 1,064,874 0.8025 2.5.0 1,311,100 0 0.0000 2.5.0 1,211,359 1064,876 0.0819 2.5.0 1,211,359 1064,876 0.0819 2.5.0 1,214,359 106,876 0.0819 2.5.0 1,215,439 281,071 0.2313 2.5.0 1,215,439 281,071 0.2313 2.5.0 1,179,525 835,801 0.0000 2.5.0 1,215,439 281,071 0.7094 2.5.0 1,311,298 955,995 0.6971 2.5.0 1,311,298 955,995 0.06971 2.5.0 1,311,298 955,995 0.06971 2.5.0 1,311,298 955,995 0.06971 2.5.0 1,311,298 955,995 0.06971 2.5.0 1,311,272 856,758 85,785 0.0663 3.03 1,225,775 367,058 0.2855 30.3 1,275,694 1,320,024 1.0323	1N − 1	07/10/96	Extraction Sample		00.00	1,532,021	o	0.000	В
2.50 1,434,419 124,377 0.0867 10.0 1,312,407 315,281 0.2439 10.0 1,312,407 317,70 0.2497 25.0 1,312,407 317,70 0.2497 25.0 1,312,407 317,626 1,039,955 0.7833 25.0 1,311,100 0.7/10/96 Extraction Sample Charcoal 0.00 1,211,100 0.000 0.0000 2.50 1,247,644 102,155 0.0819 10.0 1,215,439 281,071 0.2314 10.0 1,215,439 281,071 0.2313 25.0 1,311,298 955,995 0.6971 25.0 1,311,298 955,995 0.6971 0.2000 25.0 1,311,298 955,995 0.6971 0.3000 25.0 1,311,298 955,995 0.6971 0.3000 25.0 1,311,298 955,995 0.6091 1.311,298 955,995 0.0000 25.0 1,311,298 955,995 0.0000 25.0 1,311,298,775 85,495 0.0663 10.1 1,286,578 85,708 0.2855 30.3 1,278,694 1,320,024 1.320,024 1.320,024 1.320,024					2.50	1,393,392	122,449	0.0879	-35
10.0 1,374,428 335,281 0 10.0 1,312,407 327,737 0 25.0 1,323,184 1,061,874 0 25.0 1,323,184 1,061,874 0 0 0 1,323,184 1,061,874 0 0 0 0 1,223,184 1,061,874 0 0 0 0 0 0 0 1,223,184 1,061,874 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0					2.50	1,434,419	124,377	0.0867	5
10.0 1,312,407 327,737 0 25.0 1,323,184 1,061,874 0 25.0 1,323,184 1,061,874 0 25.0 1,327,626 1,039,955 0 2.50 1,231,359 106,876 0 2.50 1,247,644 102,155 0 10.0 1,247,644 102,155 0 10.0 1,247,644 102,155 0 10.0 1,240,262 290,664 0 10.0 1,21,319 281,071 0 25.0 1,179,525 836,801 0 25.0 1,371,298 955,995 0 1,07/10/96 Extraction Sample Process Blank 0.00 1,259,030 0 1,289,030 0 1,289,030 0 1,289,735 858,465 0 1,01 1,301,928 24,928 0 3.03 1,286,578 85,285 0 10.1 1,301,928 775 367,058 0					10.0	1,374,428	335,281	0.2439	
25.0 1,327,626 1,039,955 0 07/10/96 Extraction Sample Charcoal 0.00 1,311,100 0 0 2.50 1,231,359 106,876 0 2.50 1,247,644 102,155 0 10.0 1,247,644 102,155 0 10.0 1,247,644 102,155 0 10.0 1,247,644 102,155 0 10.0 1,247,644 102,155 0 10.1 1,21,296 280,664 0 25.0 1,179,525 836,901 0 25.0 1,371,298 955,995 0 1 07/10/96 Extraction Sample Process Blank 0.00 1,259,030 0 25.0 1,172,221 858,465 0 3.03 1,286,578 85,285 0 10.1 1,286,578 85,285 0 10.1 1,286,578 130,024 1					10.0	1,312,407	327,737	0.2497	
07/10/96 Extraction Sample Charcoal 0.00 1,311,100 0 0 0 2.50 1,231,359 106,876 0 2.50 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 102,155 0 1,247,644 10,210 0 0 1,247,255 836,801 0 0 0 1,215,439 24,955 0 1,317,298 955,995 0 1,317,298 955,995 0 1,317,296 954,958 0 1,286,578 85,285 0 1,218,694 1,320,024 1 1,310,928 1,320,024 1					25.0 25.0	1,323,184	1,061,874 1,039,955	0.8025	
2.50 1,231,359 106,876 0 2.50 1,247,644 102,155 0 10.0 1,240,262 290,664 0 10.0 1,215,439 281,071 0 25.0 1,179,525 836,801 0 25.0 1,179,525 836,801 0 25.0 1,179,525 836,995 0 1,371,298 955,995 0 25.0 1,172,221 858,465 0 25.0 1,172,221 858,465 0 1,172,221 858,465 0 1,172,221 85,775 85,775 367,058 0 10.1 1,285,775 367,058 0 1,285,775 1,286,775 1,320,024 1	1N-1	07/10/96	Extraction Sample		0.00	1,311,100	0	0.0000	
2.50 1,247,644 102,155 0 1,00,262 290,664 10.00 1,215,439 281,071 0 25.0 1,179,525 836,801 0 25.0 1,179,525 836,801 0 25.0 1,179,525 836,995 0 0 0 0 0 1,259,030 858,995 0 0 0 25.0 1,172,221 858,465 0 0 0 25.0 1,172,221 858,465 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		•			2.50	1,231,359	106,876	0.0868	
10.0 1,240,262 290,664 0 10.0 1,215,439 281,071 0 25.0 1,179,525 836,801 0 25.0 1,179,525 836,801 0 25.0 1,171,298 955,995 0 25.0 1,172,221 858,465 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0					2.50	1,247,644	102,155	0.0819	
10.0 1,215,439 281,071 0 25.0 1,179,525 836,801 0 25.0 1,179,525 836,801 0 0 0 0 1,371,298 955,995 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0					10.0	1,240,262	290,664	0.2344	
25.0 1,179,525 836,801 0 25.0 1,371,298 955,995 0 07/10/96 Extraction Sample Process Blank 0.00 1,259,030 0 0 07/10/96 Calibration Std 1.01 1,301,928 24,928 0 3.03 1,286,578 85,285 0 10.1 1,286,578 367,058 0					10.0	1,215,439	281,071	0.2313	
25.0 1,371,298 955,995 0 07/10/96 Extraction Sample Process Blank 0.00 1,259,030 0 0 07/10/96 Calibration Std 1.01 1,301,928 24,928 0 3.03 1,286,578 85,285 0 10.1 1,286,775 367,058 0					25.0	1,179,525	836,801	0.7094	
07/10/96 Extraction Sample Process Blank 0.00 1,259,030 0 25.0 1,172,221 858,465 1.01 1,301,928 24,928 3.03 1,286,578 85,285 10.1 1,285,775 367,058 30.3 1,278,694 1,320,024					25.0	1,371,298	982,995	0.6971	
25.0 1,172,221 858,465 07/10/96 Calibration Std 1.01 1,301,928 24,928 3.03 1,286,578 85,285 10.1 1,285,775 367,058 30.3 1,278,694 1,320,024	HN-1	07/10/96	Sampl			1,259,030	0	0.0000	
07/10/96 Calibration Std 1.01 1,301,928 24,928 3.03 1,286,578 85,285 10.1 1,285,775 367,058 30.3 1,278,694 1,320,024					25.0	1,172,221	858,465	0.7323	
3.03 1,286,578 85,285 10.1 1,285,775 367,058 30.3 1,278,694 1,320,024	HN-1	01/10/96	Calibration Std		1.01	1,301,928	24,928	0.0191	
1,278,694 1,320,024					3.03	1,286,578	85,285	0.0663	
1,278,694 1,320,024					10.1	1,285,775	367,058	0.2855	
				•	30,3	1,278,694	1,320,024	1.0323	

TABLE 8. (Continued)

		-		Actual cond	Tatour		
	Injection			iolloo i maani	THEFT		
Compound	Date	San Sleam S		of spike or	standard		Response
		Sampte Name	Waste Stream	std (ppm)	target area	Target area	ratio
	07/11/96	Calibration Std		1.01	1,233.886	17.241	2020.0
				3.03	1,290,644	112,816	0.0974
				10.1	1,342,405	430,339	0.3206
				30.3	1,313,941	1,395,886	1.0624
				101	1,420,602	4,829,203	3.3994
	07/11/96	Extraction Sample	Blue	0.00	1.411 591	ć	6
				2.50	1,395,047	90 481	0.0000
				2.50	1,353,532	74.111	0.0548
				10.0	1,323,621	249,112	0.1882
				10.0	1,309,466	222,575	0021
				25.0	1,316,562	745,966	0.11.0
				25.0	1,276,074	704,609	0.5522
	95/11/20	10000				•	2 1 1 1
	01/11/10	Extraction sample	Red	00.00	1,245,975	0	0.0000
				2.50	1,112,913	97,146	0.0873
				2.50	1,142,971	99,191	0.0868
				10.0	1,147,977	270,555	0.2357
				10.0	1,131,470	271,787	0.2402
				S	1,143,115	856,216	0.7490
				25.0	1,056,520	788,226	0.7461
	96/11/20	Extraction Sample	Charcoal	0.00	1 082 582	•	•
				0 4 6	796'780'7	0	0.0000
				2.30	1,052,507	98,663	0,0937
				7.50	1,001,573	87,095	0.0870
				10.0	1,009,983	213,022	0.2109
				10.0	939,402	203,360	0.2165
				25.0	975,486	702,250	0.7199
				25.0	974,223	650,953	0.6682
	07/11/96	Extraction Sample	Proсевв Blank		962,527	C	0
				25.0	886,001	610,223	0.6887
	07/11/96	Calibration Std		•			•
	•			1.01	958,729	26,447	0.0276
				3.03	970,071	61,107	0.0630
				10.1	941,918	279,091	0.2963
				30.3	931,694	894,524	0 9601

TABLE 8. (Continued)

																:	B-	.37	7																					
Response	14110	0.0057	0.0174	0.0607	0.1968	0.6278		0.0753	0.0862	7000.0	7560.0	0.1160	0.1160	0.2110	0.2051	0	0.000	0.0148	0.0146	0.0703	0.0704	0.1708	0.1795	•	0.0000	0.0065	0.0061	0.0628	0.0877	0 1875	0.101.0	0.1885	0	0.000	0.1638	1	0.0024	0.0110	0.0498	0.1/85
Target area		7,050	21,839	76,827	249,907	855,844	•	106,842	113,458	131,142	183.525	149,529	220,000	269 362	700//02	c	0 0 0	10,280	15,740	80,354	77,406	189,123	189,235		0	7,215	095'9	67,948	97,109	206,404	201 606	500,102	c	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	1751711	300.0	12 700	57.652	206.206	744,032
Internal Btandard target area		1,229,757	1,253,857	1,265,599	1,269,552	1,363,286		1,419,824	1,316,665	1,315,639	1,347,568	1,288,883	1,280,120	1,313,609	•	1,206,279	1,099,030	1.077 885	100,000	1,142,99/	1,099,497	1,107,504	1,054,513		1,1/5,556	1,107,506	1,075,616	1,081,556	1,107,330	1,100,883	1,069,669		1,107,402	1.053.244		1,146,758	1,150,405	1,158,131	1,155,481	1,217,546
Actual conc. of spike or std (ppm)		0.96	2.88	9.64	28.8	96.4	ć	0.00	2.50	2.50	10.0	10.0	25.0	25.0		00.00	2.50	2.50	10.0	0 0	0.04	0.52	75.0	00 0	2.50	00.4	00.2	10.0	10.0	25.0	25.0		0.00	25.0		96.0	2.88	9,64	28.8	96.4
Waste Stream							Rlue									Red								Charcoal	!								Process Blank				•			
Sample Name	Calibration Std						Extraction Sample	•								Extraction Sample								Extraction Sample								7	Excraction sample			Calibration Std				
Injection Date	96/80/10						96/80/10								39/80/70	06/00/10								96/80/10								96/80/20	ne lander		207.007.00	96/90/10				
Compound	CHI	2	.PI	pe	nč	li:	æ :	В							≘	41	L							GH.								Q11			21	Ē				

TABLE 8. (Continued)

												•	י כ	0																		
đi.													3-3							_				_								
Response ratio	0.0050	0.0150	0.1888	0.6122	0.0457	0.1021	0.0893	0.1194	0.1078	0.1999	0.1973	0.0000	0.0165	0.0153	0.0759	0.0785	0.1831	0.1789	0.0000	0.0116	0.0045	0.0947	0.0714	0.2050	0.2049	0.0000	0.1733	0.0023	0.0116	0.0506	0.1889	0 6366
Target area	6,589	21,729	283,807	997,162	75,369	158,665	140,762	189,536	166,444	313,058	307,060	0	22,963	21,986	104,361	103,064	242,313	237,547	0	14,313	5,616	117,391	86,793	241,822	281,017	0	203,156	2,943	14,920	62,039	241,518	982 606
incernai Btandard target area	1,311,822	1,541,279	1,502,948	1,628,767	1,650,237	1,554,143	1,576,644	1,587,228	1,544,227	1,566,114	1,556,627	1,532,021	1,393,392	1,434,419	1,374,428	1,312,407	1,323,184	1,327,626	1,311,100	1,231,359	1,247,644	1,240,262	1,215,439	1,179,525	1,371,298	1,259,030	1,172,221	1,301,928	1,286,578	1,285,775	1,278,694	1 389 700
Actual cond. of spike or std (ppm)	0.96	9.64	28.8	96.4	0.00	2.50	2.50	10.0	10.0	25.0	25.0	00.00	2.50	2.50	10.0	10.0	25.0	25.0	00.00	2.50	2.50	10.0	10.0	25.0	25.0		25.0	96.0	2.88	9.64	28.8	V. 96
Waste Stream					Blue							Red							Charcoal							Process Blank						
Sample Name	Calibration Std				Extraction Sample							Extraction Sample							Extraction Sample							Extraction Sample		Calibration Std				
Injection Date	07/10/96				07/10/96							07/10/96							01/10/96							07/10/96		07/10/96				
Compound	НО				QH							HD							GH							НО		QH				
					ĸ B								4																			

TABLE 8. (Continued)

			B-39			
Response ratio	0.0046 0.0167 0.0615 0.1997 0.6658	0.0738 0.0905 0.0921 0.1353 0.1229 0.2227	0.0000 0.0169 0.0163 0.0774 0.0789 0.1834	0.0000 0.0123 0.0123 0.0655 0.0661 0.1908	0.0000	0.0023 0.0096 0.0474 0.1815
Target area	5,696 21,522 82,569 262,380 945,813	104,123 126,282 124,626 179,075 160,970 293,165	18,817 18,651 88,840 89,223 209,593	12,895 12,299 66,175 62,123 186,163	0 151,386	2,161 9,277 44,600 169,133 689,468
Internal standard target area	1,233,886 1,290,644 1,342,405 1,313,941 1,420,602	1,411,591 1,395,047 1,353,532 1,323,621 1,309,466 1,316,562	1,245,975 1,112,913 1,142,971 1,147,977 1,131,470 1,143,115 1,056,520	1,082,582 1,052,507 1,001,573 1,009,983 939,402 975,486	962,527 886,001	958,729 970,071 941,918 931,694 1,071,736
Actual conc. of spike or std (ppm)	0.96 2.88 9.64 28.8 96.4	0.00 2.50 2.50 10.0 25.0	0.00 2.50 2.50 10.0 25.0	0.00 2.50 2.50 10.0 10.0 25.0	k 0.00 25.0	0.96 2.88 9.64 28.8
Waste Stream		Blue	Red	Charcoal	Procевв Blank	
Sample Name	Calibration Std	Extraction Sample	Extraction Sample	Extraction Sample	Extraction Sample	Calibration Std
Injection Date	07/11/96	07/11/96	07/11/96	07/11/96	07/11/96	07/11/96
Compound	을 Appendix	QH X B	≘ 143	QI		GII

TABLE 8. (Continued)

																		B-	40																		
		Reaponse	ratio	0.0032	0.0101	0.0552	0.2427	1.0164	0,0000	0.0017	0.0021	0.0047	0.0063	0.0107	0.0145	0.1877	0,2305	0.2564	0.3153	0.3949	0.3998	0.4973	0.2513	0.2897	0,3120	0.3638	0.3555	0.4765	0.5454	0.000	0.3039	6	0.0009	0.0036	0.0265	0.1780	0.9333
			Target area	3,919	12,645	69,903	308,109	1,385,695	0	2,197	2,737	6,378	8,158	13,684	18,993	226,406	253,290	276,356	360,404	434,218	442,814	524,429	295,461	320,802	335,573	393,428	393,603	524,548	583,422	0	320,081	,	1,046	4,141	30,670	205,640	1,136,327
	Internal	standard	target area	1,229,757	1,253,857	1,265,599	1,269,552	1,363,286	1,419,824	1,316,665	1,315,639	1,347,568	1,288,883	1,280,120	1,313,609	1,206,279	1,099,030	1,077,885	1,142,997	1,099,497	1,107,504	1,054,513	1,175,556	1,107,506	1,075,616	1,081,556	1,107,330	1,100,883	1,069,669	1,107,402	1,053,244	1 146 760	BC/ 1011/1	1,150,405	1,158,131	1,155,481	1,217,546
(penu	Actual conc.	of apike or	std (ppm)	0.81	2.44	8,12	24.4	81.2	0.00	5.00	5.00	10.0	10.0	25.0	25.0	0.00	5.00	5.00	10.0	10.0	5	25.0	0.00	5.00	5.00	10.0	10.0	25.0	25.0		25.0	נמיי		2.44	8.12	24.4	81.2
(Continued)			Waste Stream						Blue							Red							Charcoal							Process Blank					٠		
		,	Sample Name	Calibration Std					Extraction Sample							Extraction Sample							Extraction Sample							Extraction Sampie		Calibration Std					
		Injection	Date	03/80/10					07/08/53							07/08/95							96/80/10							96/80/10		96/80/10	•				
		í	Compound	L-Der	Αŗ	pe	en	diz	R L-Der							L-Der	3	L 4	4				L-Der							L-Der		L-Der					

TABLE 8. (Continued)

	•						•	
Compound	Date	Sample Name	Waste Stream	std (ppm)	target area	Target area	ratio	
L-Der	07/10/96	Calibration Std		0.81	1,311,822	2,167	0.0017	
A				2.44	1,452,174	9,186	0.0063	
PF				8.12	1,541,279	66,386	0.0431	
eı				24.4	1,502,948	346,988	0.2309	
ndi				81.2	1,628,767	1,645,193	1.0101	
X L-Der	07/10/96	Extraction Sample	Blue	00.00	1,650,237	0	0,000	
В				5.00	1,554,143	1,159	0.0007	
				5.00	1,576,644	956	0.0006	
				10.0	1,587,228	4,507	0.0028	
				10.0	1,544,227	3,760	0.0024	
				25.0	1,566,114	11,164	0.0071	
				25.0	1,556,627	8,270	0,0053	
L-Der	07/10/96	Extraction Sample	Red	0.00	1,532,021	328.115	0.2142	E
1				5.00	1,393,392	364,171	0.2614	3-4
45				5.00	1,434,419	440,722	0.3072	1
				10.0	1,374,428	466,507	0.3394	
				10.0	1,312,407	449,077	0.3422	
				25.0	1,323,184	772,965	0.5842	
				25.0	1,327,626	682,025	0.5137	
							,	
L-Der	07/10/96	Extraction Sample	Charcoal	00.00	1,311,100	100,708	0.0768	
				5.00	1,231,359	199,744	0.1622	
				5.00	1,247,644	80,075	0.0642	
				10.0	1,240,262	165,545	0.1335	
				10.0	1,215,439	221,708	0.1824	
				25.0	1,179,525	373,331	0.3165	
				25.0	1,371,298	621,268	0.4531	
L-Der	07/10/96	Extraction Sample	Process Blank		1,259,030	0	0.000	
				25.0	1,172,221	384,564	0.3281	
L-Der	01/10/96	Calibration Std		0.81	1,301,928	1.313	0100	
				2.44	1.286.578	240/2	0100.0	
				8.12	1,285,775	004.00	0.000	
				1 7 7 7	1 220 604	676,66	0.0307	
		•		7 (1,2/8/84	252,444	0.1974	
				91.2	1,386,790	1,349,320	0.9716	

TABLE 8. (Continued)

	100,40			Actual conc.	Internal			
to a code	Dato			of spike or	standard		Response	
compound	Date	Sample Name	Wagte Stream	std (ppm)	target area	Target area	ratio	
L-Der	07/11/96	Calibration Std		0.81	1,233,886	2,519	0.0020	
				2.44	1,290,644	10,491	0.0081	
				8.12	1,342,405	65,315	0.0487	
				24.4	1,313,941	301,411	0.2294	
				81.2	1,420,602	1,436,699	1.0113	
g L-Der	07/11/96	Extraction Sample	Blue	00.00	1,411,591	0	0,0000	
				5.00	1,395,047	3,775	0,0027	
				5.00	1,353,532	4,981	0.0037	
				10.0	1,323,621	11,229	0.0085	
				10.0	1,309,466	11,816	0.0000	
				25.0	1,316,562	32,910	0.0250	
				25.0	1,276,074	25,252	0.0198	
L-Der	07/11/96	Extraction Sample	Red	00.00	1,245,975	258,142	0.2072	I
				5.00	1,112,913	328,379	0.2951	3-4
				5.00	1,142,971	370,588	0.3242	12
				10.0	1,147,977	287,140	0.2501	
				10.0	1,131,470	369,789	0.3268	
				25.0	1,143,115	685,787	0.5999	
				25.0	1,056,520	509,543	0.4823	
L-Der	07/11/96	Extraction Sample	Charcoal	0.00	1,082,582	145,636	0.1345	
				5.00	1,052,507	134,418	0.1277	
				2.00	1,001,573	219,399	0.2191	
				10.0	1,009,983	240,968	0.2386	
				10.0	939,402	206,450	0.2198	
				25.0	975,486	508,266	0.5210	
				25.0	974,223	349,760	0.3590	
L-Der	96/11/60	Extraction Sample	Process Blank		962,527	0	0.0000	
			•	25.0	886,001	256,091	0.2890	
L-Der	07/11/96	Callbration Std		0.81	958,729	0	0.0000	
				2.44	170,079	3,558	0,0037	
				8.12	941,918	19,218	0.0204	
				24.4	931,694	129,540	0.1390	
				81.2	1,071,736	948,877	0.8854	

Appendix B

RESULTS FOR CALIBRATING STANDARDS ANALYZED WITH THE WASTE STEAM SAMPLE TABLE 9.

Compound	Injection Date	Intercept	Std.Err., Intercept	Slope coefficient	Std.Err., Slope	Slope2 coefficient	Std.Err., Slope2	Average Deviation from Regression
≘ 1	07/08/96 07/10/96 07/11/96	003341 004171 006996	0.003076 0.002866 0.003524	0.00647066 0.00652606 0.00685843	0.000068 0.000063 0.000078	1 1 1	1 1 1	0.007683 0.007157 0.008801
1-NE 47	07/08/96 07/10/96 07/11/96	0.013514 008879 010830	0.042343 0.012842 0.023070	0.03149511 0.03274973 0.03287345	0.000893 0.000271 0.000487	1 1 1	I I 1	0.105714 b 0.032063 0.057599
L-Der	07/08/96 07/10/96 07/11/96	011741 014030 009691	0.015611 0.008886 0.021921	0.00750617 0.00769532 0.00606664	0.001630 0.000928 0.002289	0.000057242 0.000057693 0.000070627	0.00001900	0.030347

Regression lines or curves on each respective date were used to compute back-calculated compound concentrations (ppm) shown in Table 10.

					Mean				
Compound	Sample Name	injection Date	of apike or atd (ppm)*	z	Response Ratio	Std. Dev.	Coef. of Variation	Back-calculated conc. (ppm)	Percent Recovery
HD	Blue Wasta Stream	96/80/10	0.0	-	0.0753	1]	12.15	
A		-	2.5	7	0.0929	9600.0	10.28	14.88	109.3
pp			10.0	7	0.1261	0.0143	11.31	20.00	78.6
en			25.0	7	0.2080	0.0042	2.02	32.67	82.1
£ dix	Blue Waste Stream	07/10/96	0.0	-	0.0457**	1	1	7.64	1
: F			2.5	7	0.0957	0.0091	9.47	15,30	306.5
3			10.0	7	0.1136	0.0082	7.24	18.05	104.1
			25.0	8	0.1986	0.0019	0.94	31.07	93.7
HD	Blue Waste Stream	07/11/96	0.0	-	0.0738	ı	ı	11.78	ı
			2.5	7	0.0913	0.0011	1.20	14.33	102.3
			10.0	7	0.1291	0.0087	6.17	19,85	80.7
			25.0	7	0.2205	0.0030	1,38	33,17	85.6
≘ 1	Red Waste Stream	96/80/10	2.5	7	0.0147	0.0001	1.01	2.79	111.6
.43		•	10.0	7	0.0704	0.0001	0.10	11,39	113.9
В			25.0	7	0.1751	0.0061	3.51	27.58	110.3
EFD.	Red Waste Stream	07/10/96	2.5	2	0.0159	0.0008	5.12	3.08	123.0
			10.0	7	0.0772	0.0018	2.38	12.47	124.7
			25.0	7	0.1810	0.0030	1.64	28.38	113.5
GH	Red Waste Stream	07/11/96	2.5	2	0.0166	0.0004	2.51	3.44	137.7
			10.0	7	0.0781	0.0010	1.33	12.41	124.1
			25.0	7	0.1813	0.0030	1,63	27.45	109.8
IID	Charcoal Waste Stream	07/08/96	2.5	7	0.0063	0.0003	4.66	1.49	59.6
			10.0	7	0.0753	0.0176	23.37	12.15	121.5
			25.0	7	0.1880	0.0007	0.37	29.57	118.3
GH	Charcoal Waste Stream	n 07/10/96	2.5		0.0081	0,0050	62.47	1.87	75.0
			10.0	7	0.0830	0.0164	19.79	13,36	133.6
			25.0	7	0.2050	0.0001	0.03	32,05	128.2
GII	Charcoal Waste Stream : 07/11/96	96/11/20 w	2.5	7	0.0123	0.0000	0.16	2.81	112.3
			10.0	7	0.0658	0.0004	0.65	10.62	106.2

TABLE 10. (Continued)

Compound	Sample Name	Injection Date	Actual conc, of spike or std (ppm)	Mean Response N Ratio	Std. Dev.	Coef. of Variation	Back-calculated	Percent Recovery
E Appendix	Calibration Standard	01/08/96	0.96 2.88 9.64 28.80 96.40	2 0.0041 2 0.0142 2 0.0552 2 0.1877 2 0.6194	0.0024 0.0045 0.0077 0.0130	58.52 31.65 13.98 6.91	1.14 2.72 9.05	
GE B	Calibration Standard	07/10/96	0.96 2.88 9.64	2 0.0036 2 0.0133 2 0.0545		53.64 17.93	96.25 1.20 2.67	1 1 1
GII	Calibration Standard	07/11/96	28.80 96.40	2 0.1889		0.02	9.00 29.58 96.24	1 1 1
149		.	<i>₹</i> .	2 0.0034 2 0.0131 2 0.0544 2 0.1906 2 0.546	0.0017 0.0050 0.0100 0.0128 0.0159	48.63 38.33 18.39 6.74 2.43	1.52 2.93 8.96 28.81	B-45
GH	Process Blank	07/08/96 07/10/96 07/11/96	25.00 25.00 25.00	1 0.1638 1 0.1733 1 0.1709	1 1 1	111	25.83 27.20 25.93	103.3 108.8 103.7

TABLE 10. (Continued)

Compound	Sample Name	Injection Date	Actual conc. of spike or std (ppm)	z	Mean Response Ratio	Std. Dev.	Coef. of Variation	Back-calculated	Percent
T E Appendi	Calibration Standard	07/08/96	1.01 3.03 10.10 30.30	2222	0.0265 0.0823 0.3253 1.0395	0.0093 0.0225 0.0623 0.1049	35.26 27.30 19.16 10.09	0.41 2.18 9.90 32.58	RECOVERY
х В - Т-	Calibration standard	07/10/96	1.01		0.0233 0.0755 0.2980	0.2463	7.76 25.38 17.18	100.37 0.98 2.58	1 1 1
150	Calibration Standard	07/11/96	30.30 101.00 1.01 3.03 10.10	ପପ ପଠାଁଠ	1,0361 3,2859 0,0289 0,0752	0.0053 0.0147 0.0018 0.0173	5.93 0.51 0.45 6.36 22.96	9.37 31.91 100.60 1.21 2.62	B-4
HN-1	Process Blank	07/08/96 07/10/96 07/11/96	30.30 101.00 25.00 25.00 25.00	11 22	1,0112 3,3032 0,6694 0,7323 0,6887	0.1360	7.15	9.71 31.09 100.81 20.82 22.63 21.28	89.5 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

TABLE 10. (Continued)

Compound	Sample Name	Injection Date	<pre>hctual conc. of spike or std (ppm)</pre>	z	Mean Response Ratio	Std.	Coef. of	Back-calculated	Percent
HN-1	Blue Waste Stream	07/08/96		,		,	Hotant in	(bbill)	кесочегу
		no Inn I in	C * 7	V	0.0433	0.0018	4.20	0.94	37.8
			10.0	7	0.1221	0.0009	0.76	3.45	3.4.5
enc			25.0	7	0.4290	0.0168	3.92	13.19	52.8
1 2 2	Blue Charles Charles	707077	(,	,				
T K	biue wastu stream	01/10/96	2.5	~	0.0522	0.0023	4.33	1.86	74.6
			10.0	7	0.1463	0.0026	1.77	4.74	47.4
			25.0	7	0.5137	0.0164	3.20	15,96	63.8
HN-1	Blue Waste Stream	01/11/96	2.5	~	0.0562	1,000	6		
			10.0	c	1921 0	44000	50.0	7.04	81.6
			25.0	٠,	0.5594	0.0129	7.20	5.78	57.8
) • !			2010.0	1.82	17,35	69.4
1: 1:	Red Waste Stream	96/80/10	2.5	7	0.0813	0.0017	2.13	2.15	. 90
			10.0	7	0.2135	0.0006	0,30	6.35	63.5
			25.0	7	~ 0.7368	0.0243	3.29	22,96	91.6
,			,						•
T - NII	Ked Waste Stream	01/10/96	2.5	7	0.0873	0.0008	0.95	2.94	117.5
			10.0	7	0.2468	0.0041	1.66	7.81	נ מר
			25.0	7	0.7929	0.0136	1.71	24.48	97.9
20									
7 - NII	Red waste Stream	07/11/96	2.5	7	0.0870	0.0004	0.41	2.98	119.1
			10.0	~	0.2379	0.0032	1.35	7.57	75.7
			25.0	7	0.7475	0.0021	0.28	23.07	92.3
HN-1	Charcoal Waste Stream	96/80/10	2.5	7	0.0862	0.0103	11 94	ć	- 1
			10.0	7	0.1984	00100	FC 11	16.2	92.3
			25.0	c	0009	0010.0	#O.C	78.5	58,7
			1	4	0.0000	0.0258	3.75	21.45	85.8
HN-1	Charcoal Waste Stream	07/10/96	2,5	7	0.0843	0.0035	4.12	2 85	6
			10.0	7	0.2328	0.0022	ν ₀ Ο		6.611
			25.0	7	0.7033	0.0087	1.24	21.75	/3.8 87.0
1 - NE	Charcoal Waste Stream	07/11/96	20.00	~	0.0903	0.0048	5.31	3.08	123.1
			10.0	7	0.2137	0.0039	1.84	6.83	68.3
			2	c					

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	Percent	Recovery	35.8	22.6	12.7	37.		10.3	2 (4	0.90	20.0	i	107.1	152.6	91.	1	128.1	112.8	110.1	ı	193.6	77.8	112.	•	89.5	95.4	86,	1	79.2	85.9	115.9	i	85.3	99.5	113.1
	Back-calculated	conc. (ppm)	1.79	۲.	3.16	1.88	2.13	2.58	2.07	2.94	5.00	22.65	Θ	37.91	45.52	24.98	31,38	36,26	52.50	27.16		34.94	55.25	28.75	33,22	38.29	50,32	10.91	4	19.50	39.89	19.39		29,35	47.67
	Coef. of	Variation	15.53	20.41	21.20	14.65	10.85	20.64	21.57	4.36	16.45		7.53	15.85	15.37	ī	11.41	0.57	9.08	. 1	99.9	18,80	15.37	1	5.25	1.63	9.54		61.24	21.91	25.09	ı	37.25	5.81	26.04
	std.	Dev.	0.0003	0.0011	0.0027	0.0001	0.0003	0.0013	0.0007	0.0004	0.0037	ı	0.0183	0.0563	0.0689	1	0.0325	0.0020	0.0498	ı	0.0206	0.0542	0.0832	ι	0.0158	0.0059	0.0488	I	0.0693	0.0346	0.0965	ı	0.0646	0.0133	0.1146
Mean	Rевропве	Ratio	0.0019	0,0055	0.0126	0.0007	0.0026	0.0062	0.0032	0.0088	0.0224	0.1877	0.2434	0,3551	0.4486	0.2142	0.2843		0.5489	0.2072	0.3096	0.2885	0.5411	0.2513	0.3008	•	0.5110	0.0768	0.1132	0.1579	0.3848	0.1345	0.1734	0.2292	0.4400
		z	7	7	7	2	2	2	7	7	~	-	2	7	7	-	2	2	7	-	2	2	7	7	7	7	7	-	7	7	7	ન	7	7	2
Actual conc.	of apike or	erd (ppm)	2	10	25	ហ	10	25	ហ	10	25	0	ស	10	25	0	S	10	25	0	'n	10	25	0	2	10	25	0	S	10	25	0	ភ	10	25
-	Injection	расе	96/80/10			96/01/10	•		07/11/96	•		07/08/96				01/10/96				07/11/96	•			. 96/80/10		•		07/10/96				07/11/96			
		Sample Name	Blue Waste Stream			Blue Wastn Stream			Blue Waste Stream			Red Waste Stream				Red Waste Stream				Red Waste Stream				Charcoal Waste Stream				Charcoal Waste Stream				Charcoal Waste Stream			
	Pario Cano C	compound	A L-Der	.pı	pen	p. L-Der	x	В	L-Der			L-Der				L-Der	52	2		L-Der				L-Der				L-Der				L-Der			

Cempound	Sample Name	Injection Date	Actual conc. of splke or std (ppm)	Z	Mean Response Ratio	Std. Dev.	Coef, of Variation	Back-calculated	Percent
Ap l-Der	Calibration Standard	96/80/10	0.81	7	0.0020	0.0016	78.48	(Ppm)	recovery
pen			2.44	2 2	0.0068	0.0046	67.02	2.43	1 1
ndix			24.40 81.20	1 7 7	0.2103	0.0203 0.0458 0.0588	49.76 21.76 6.03	6.67	i I
WL-Der	Calibration Standard	96/01/10	0.81	7	0.0013	0.0005		81.18	I
			2.44 8.12	2 2	0.0057	0.0009	15.98	1.97	1 1
			24.40 81.20	2 2	0.2141 0.9908	0.0237	23.62	6.32 24.98	1 1
L-Der	Calibration Standard	07/11/96	0.81	~	0100	2120.0	2.75	81.18	ı
			2.44	. 67 (0.0059	0.0032	141.42 53.48	1.73	t
153			24.40 81.20	7 7 7	0.0345 0.1842 0.9483	0.0200 0.0639 0.0891	57.86 34.68 9.39	5.76 24.80	1 1 1
L-Der	Process Blank	07/08/96 07/10/96 07/11/96	25.00 25.00 25.00		0.3039	1 1	1 1	33.50	B-49 0.981
				,	0.2890	1	1	34.99	140.0

When no agent was detected at the O ppm level(no spike)in the waste stream, the O ppm level is not included in this

Value rejected based on the Q test. GC injection problem for this sample. Therefore, the HD results for this day are not included in the average values reported in Table 11.

TABLE 11. SUMMARY OF DESCRIPTIVE STATISTICS FOR WASTE STREAM SAMPLES OVER THREE DAYS OF ANALYSIS

				B-	·50		
Std.Dev., Percent Recovery	t 1 1	1 1	3.0		34.5 14.8 2.7	12.0 5.6 2.7	27.9 20.6 6.0
Percent Recovery	1 1 1	I I	105.3	1	105.8 79.6 83.8	124.1 120.9 111.2	82.3 120.4 120.7
(ppm) c.v.	23.3	1.8	2.9	2.2	6.3 7.4 1.6	9.7 4.7 2.4	33.9 17.1 5.0
Back-calculated Conc. Mean Std.Dev.	0.30	1.28	0.76	0.26	0.91 1.47 0.54	0.30 0.56 0.68	0.70 2.06 1.49
Back-calcu Mean	1.29 2.77 9.00	29.30 96.31	26.32	11.96	14.60 19.92 32.92	3.10 12.09 27.80	2.06 12.04 30.16
Mean Response Ratio	0.0037 0.0135 0.0547	0.1890	0.1693	0.0745	0.1276 0.2143	0.0157 0.0752 0.1791	0.0089 0.0747 0.1947
z	9 9	9	m	~	7 4 4	و بې و	999
Actual conc. of spike or std (ppm)*	0.96 2.88 9.64	28.80 96.40	25.00	0.00	10.00	2.50 10.00 25.00	2.50 10.00 25.00
Sample Name	Calibration Standard		Procese Blank	Blue Waste Stream		Red Waste Stream	Charcoal Waste Stream
Compound	≘ Appen		QH	≘ 154	4	П	НО

TABLE 11. (Continued)

					B- 51	
Std.Dev. Percent Recovery		1 1	3.8	21.1	16.7	15.6 7.0 2.6
Percent		F 1	86.3	64.6 46.5 62.0	107.5 72.4 94.0	109.8 66.9 86.2
(ppm) C.V.	46.0	6.0	4.4	32.7 22.7 12.4	15.5 9.7 3.6	14.2 10.5 3.0
Back-calculated Conc. Mean Std.Dev.	0.40 0.48 0.98	1.91 3.97	0.94	0.53 1.06 1.92	0.42 0.70 0.85	0.39 0.70 0.65
Back-calcı Mean	0.87 2.46 9.66	31.86 100.60	21.58	1,62 4,65 15,50	2.69 7.24 23.51	2.74 6.69 21.55
Mean Response Ratio	0.0262 0.0776 0.3106	3.2546	0.6968	0.0505 0.1492 0.5007	0.0852 0.2328 0.7591	0.0870 0.2150 0.6955
z	999	9 9	m	9 9 9	9 9 9	و بو ی
Actual conc. of spike or std (ppm)	1.01 3.03 10.10	101.00	25.00	2.50 10.00 25.00	2.50 10.00 25.00	2.50 10.00 25.00
Sample Name	Calibration Standard		Process Blank	Blue Waste Stream	Red Wasts Stream	Charcoal Waste Stream
Compound	I E Appendi	ix B		HN-1	155	111111111111111111111111111111111111111

TABLE 11. (Continued)

Compound	Sample Name	Actual conc. of spike or std (ppm)	z	Mean Response Ratio	Back-calcu	Back-calculated Conc.	(wdd)	Percent	Std.Dev., Percent
							;	vecuvet y	vecover y
L-Der	Calibration Standard	0.81	9	0.0015	1.84	0.18	9.6	ı	1
		2.44	9	0.0061	2,48	0.35	14.1	1	ı
		8.12	9	0.0374	6.56	1.76	26.8	1	ı
		24.40	9	0.2029	24.80	3.73	15.0	ı	ı
		8.1.20	9	0.9713	81.16	2.85	3.5	ı	1
L-Der	Process Blank	25.00	æ	0.3070	34.55	0.92	2.7	138.2	3.7
L-Der	Blue Waste Stream	5.00	9	0.0019	1.92	0.14	7.3	38.3	2,8
		10.00	9	0.0056	2.44	0.39	16.1	24.4	3.9
		25.00	9	0.0137	3.58	1.17	32.5	14.3	4.7
L-Der	Red Waste Stream	00.0	_. m	0.2030	24.93	2.25	9.0	1	i
		5.00	9	0.2791	32.06	4.33	13.5	142.7	52.8
		10.00	9	0.3281	36.32	3,35	9.5	113.9	45.5
		25.00	9	0.5129	51.03	5,98	11.7	104.4	18.9
L-Der	Charcoal Waste Stream	00.00	က	0.1542	19.68	8.92	45.3	ı	1
		5.00	9	0.1958	23.80	9.45	39.7	89.7	81.3
		10.00	9	0.2489	29.03	8.59	29.6	93.5	18.0
		25.00	9	0.4453	45.83	7.41	76.2	104 6	3 70

* When no agent was detected on the 0 ppm level (no spike) in the waste stream, the 0 ppm level is not included in this table.

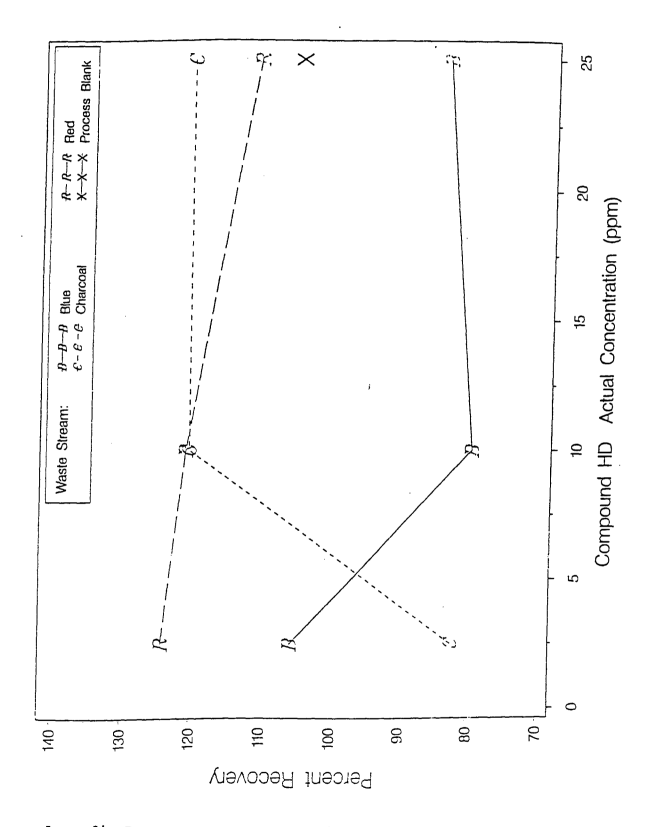


FIGURE 4. PERCENT RECOVERY OF HD FROM WASTESTREAM SAMPLES AVERAGED OVER THREE INJECTION DATES (EXCLUDING 7/10/96 FOR BLUE WASTESTREAM).

Appendix B

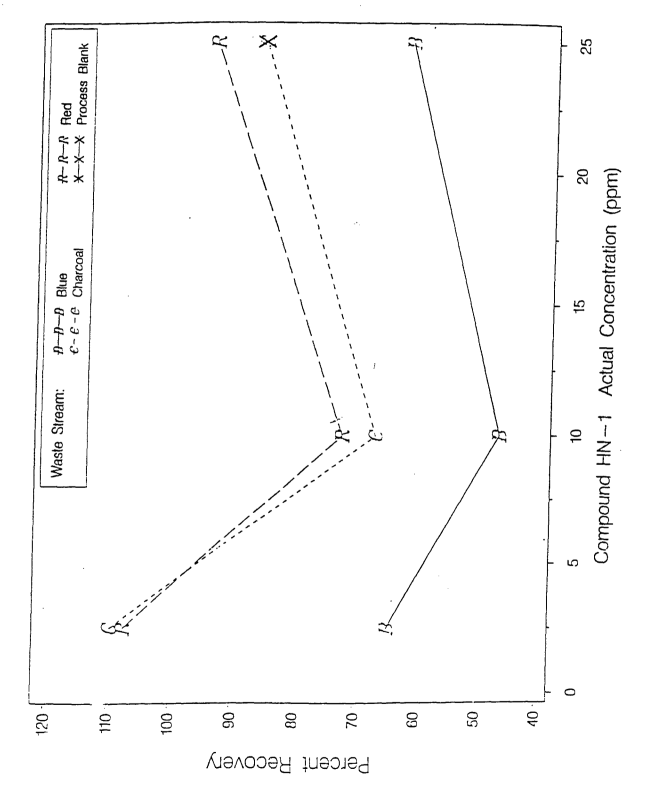


FIGURE 5. PERCENT RECOVERY OF HN·1 FROM WASTESTEAM SAMPLES AVERAGED OVER THREE INJECTION DATES (7/8-11/96).

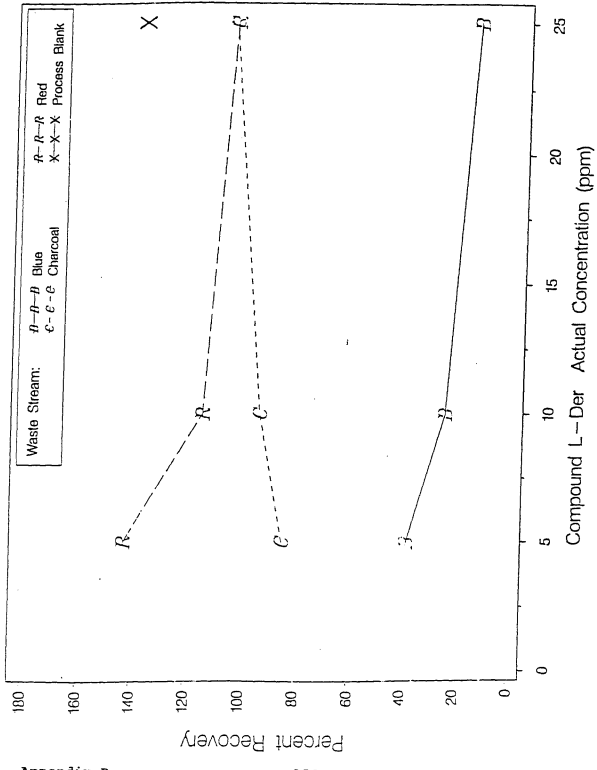


FIGURE 6. PERCENT RECOVERY OF L-Der FROM WASTESTREAM SAMPLES AVERAGED OVER THREE INJECTION DATES (7/8-11/96).

Appendix B

ATTACHMENT C

Analysis of Waste Streams

Task 95-38 Chemistry Report G155538A Summary of Waste Stream Analyses

On June 20 and 26, 1996, two waste streams (Blue Process June 17, 1996 Lot # 96-0037-057 and Red Process June 13, 1996 Lot # 96-0037-055) were analyzed by GC/MSD using the extraction procedure provided by the client on June 10, 1996. A low level of HD was found in the Blue sample and a low level of L was found in the Red sample. HN-1 was not detected in either sample, L was not detected in the Blue sample, and HD was not detected in the Red Sample. The results given in the tables were determined using regression parameters for calibration standards from 1 to 100 ppm based on a quadratic regression model. The data is recorded in LRB-356 and Task 38 Binder-02.

Table 1. Blue Waste Stream Analysis Results

Date	HD	L	HN-1
June 20	6.6 ppm	BDL	BDL
June 26	5.5 ppm	BDL	BDL
	Stal	istics	
Average	6.1 ppm		
Std Dev	0.8 ppm		

Table 2. Red Waste Stream Analysis Results

Date	HD	L	HN-I
June 20	BDL	9.3 ppm	BDL
June 26	BDL	13.2 ppm	BDL
		Statistics	
Average		11.3 ppm	
Std Dev		2.8 ppm	

Where BDL is below detection limit.

Table 3 summarizes the results for the analysis of RRS waste streams during August, 1996. The values given in the Table 3 represent the average for duplicate determinations using regression parameters for calibration standards from 1 to 100 ppm based on a quadratic regression model for L and simple linear regression models for HD and HN-1. The data are recorded in LRB-356 and Task 38 Binder-02.

Table 3. Waste Stream Analysis Results

Analysis and Dosing Date	Waste Stream*	HN-1	HD	L
August 13, 1996	Blue - 11/28/95	BDL	16.2 ppm	BDL
August 13, 1996	Red - 11/28/95	BDL	BDL	3.8 ppm ^c
August 13, 1996	Charcoal - 1/25/96	BDL	BDL	7.8 ppm ^d
August 29, 1996	Charcoal - 950096-064	BDL	BDL	74.2 ppm

^a The lot number or date received is used to identify the waste stream. ^b BDL is below the detection limit. ^c Value reported but it is below the method detection limit. ^d Value reported but the qualifiers were not satisfied and the value is below the method detection limit.

Since the qualifying ions were not satisfied for L in the Charcoal sample analyzed on August 13, full scan analyses were performed on the Charcoal - 950096-064 samples to verify the presence of L-Der. The mass spectra clearly show that the L-Der is present in the Charcoal - 950096-064 waste stream. The L value reported for Charcoal - 950096-064 (74.2 ppm) was higher than the value reported by Brian MacIver at ERDEC before shipment (10.7 ppm). This is the first analysis where the MREF measurement has been different than the value reported by ERDEC or by Battelle at Edgewood.

The relative oxidizing strength of each waste stream was determined as part of the analysis procedure. The oxidizing strength of the waste streams are compared to a fresh solution of DCDMH. The results for all of the waste streams are shown in Table 4.

Table 4. Relative Oxidizing Strength

Waste Stream*	Analysis Date	% [Ox] ^b
Charcoal - 950096-064	9/3/96	100%
Blue - 11/28/95	8/15/96	2%
Red - 11/28/95	8/15/96	7%
Charcoal- 1/25/96	8/15/96	3%
Red - 96-0037-047	6/21/96	25%
Red - 96-0037-055	6/21/96	33%
Blue - 96-0037-049	6/21/96	1%
Blue - 96-0037-057	6/21/96	1%
Charcoal - 95-0096-014L-018HN-1-023HD	6/21/96	5%
Charcoal- 1/25/96	6/21/96	3%

^a The lot number or date received is used to identify the waste stream. ^b Relative oxidizing strength expressed in percent.

APPENDIX C

Studies Performed at ERDEC

PRODUCT ANALYSIS OF "ARCHIVED" WASTESTREAM* ("Blue" Process Chemistry)

tertiary-butanol chloroform	22.8
	25.7
trichloroethene	0.1
CH_Cl	0.4
CH ₃ -C-OH	
1	
	0.2
	0.8
CH ₂ CI	0.2
CH C OH	
CH ₃ -C-OH	
CH CI	
	0.5
	0.4
	0.4
. Ω	25.0
<u>'0</u>	
•	1.2
	17.7
	3.5
	1.45
· ·	4.43
	CH ₃ -C-OH CH ₃ dichlorobutene dichlorobutene CH ₂ Cl CH ₃ -C-OH CH ₂ Cl trichlorobutane CH ₂ Cl trichlorobutane CICH=CH-S-CH ₂ CH ₂ Cl O CICH ₂ CHCl-S-CH ₂ CH ₂ Cl isomer CICH ₂ CHCl-S-CH ₂ CH ₂ Cl isomer CICH ₂ CHCl-S-CH CICH ₂ Cl isomer CICH ₂ CHCl-S-CH CICH ₂ Cl isomer CICH ₂ CHCl-S-CH CICH ₂ Cl isomer

⁽a) Studies performed at ERDEC, composition analysis conducted via the GC-MS chemical

⁽a) Studies performed at ERDEC, composition analysis conducted via the OC-IVIS chemical ionization (CI) mode.
(b) Area % calculated from the Total Ion Chromatogram (TIC) of the mass spectrometer.
(c) Area % is semi-quantative, the intent is to show the percent of the peak in comparison to other peaks in the chromatogram. Peaks less than 0.1% of the TIC are not quantitated.
(d) Method Quantitation Limit (MQL) for agent is (50 ppm).

PRODUCT ANALYSIS OF "ARCHIVED" WASTESTREAM'

an (sec)	Compound	Area % h,c,4		
24	Tertiary-butanol	13.7		
27	<u>chloroform</u>	46.7		
32	OH	1.5		
	/			
	Cl ₂ CCH			
	\ 			
	OH			
36	CH ₂ Cl	7.2		
	CH,C-OH			
	CH,			
40	Cl ₂ CHCH ₂ -OH	0.4		
53	dichlorobutene + unknown (MW = 140)	0.3		
5 6	unknown	1.3		
69	trichlorobutane	0.4		
75	CH₃C1	1.5		
	1			
	CH ₃ C-OH			
	1			
	CH <u>.</u> Cl			
81	trichlorobutene	0.1		
94	trichlorobutane	4.6		
96	trichlorobutene '	0.2		
99	trichlorobutene	0.3		
108	unknown (MW = 174)	0.05		
136	trichlorobutene	1.4		
148	unknown (MW = 208)	0.1		
155	tetrachlorobutane	0.3		
167	CH ₂ Cl	2.1		
	1			
	CICH ₂ -C-OH isomer			
	l			
	CH ₂ Cl			
188	scan 167 isomer	0.05		
205	scan 167 isomer	0.2		
220	scan 167 isomer	1.6		
234	scan 167 isomer	0.2		
292	tetrachlorobutene	0.2		
328	unknown (MW = 216)	0.03		
340	unknown (MW = 206)	0.06		
407	_1 N	15.2		
	=0			
	/, — √ — ·			
	σ "			
518	unknown (MW = 242)	0.1		
556	unknown	0.05		
510	unknown	0.04		

(a) Studies conducted at ERDEC, composition analysis conducted via the GC-MS chemical ionization (CI) mode
(b) Area % calculated from the Total Ion Chromatogram (TIC) of the mass spectrometer
(c) Area & is semi-quantative, the intent is to show the percent of the peak in comparison to other peaks in the chromatogram. Peaks less than 0.1% of the TIC are not quantitated.
(d) Method Quantitation Limit (MQL) for agent is (50 ppm).

Product Analysis of "Archived" Wastream

Scan (sec)	("Charcoal" Process Chemist Compound	Area % bc		
37	OH	25.0		
_	/			
	CI,CCH			
	\			
	OH			
39	Dichlorobutane	17.5		
60	Trichloroethylene	6.5		
79	Dichlorobutyl alcohol	1.1		
87	Trichlorobutene	1.5		
98	Dichlorobutene	7.6		
138	0	4.4		
	II			
	CICH, CH, SCI			
	11			
	0			
141	Trichlorobutene	3.3		
172	Trichlorobutene	3.4		
205	CH ₂ CH ₂ CI	3.2		
	/			
	HN			
	CTT CTT CT			
217	CH,CH,Cl Unknown	4.6		
225	Trichlorobutene	2.0		
297		0.6		
369	· · · · · · · · · · · · · · · · · · ·			
507		Not quantitated		
	NC1			
	0 101			
350-500	→ NH o	Not quantitated		
	→ ⁰			
	—NH			
483	Hexachlorobutene	15.9		
523	0	1.0		
	II · · · ·			
	CI-CH ₂ CHCI-S-CHCICH ₂ Cl isomer			
537	0	1.7		
	11			
	CI-CH ₂ CH ₂ -S-CHClCH ₂ Cl isomer			
	11			
566	0	0.4		
566	0	V. 4		
	CI-CH,CHCI-S-CHCICH,CI isomer			
	 0			
579	0	0.2		
4.7	11	v		
	CI-CH ₂ CHCI-S-CHCICH ₂ CI isomer			

⁽a) Studies performed at ERDEC, composition analysis conducted via the ge-ms chemical ionization (CI) mode
(b) Area % calculated from the Total Ion Chromatogram (TIC) of the mass spectrometer.
(c) Area % is semi-quantative, the intent is to show the percent of the peak in companison to other peaks in the chromatogram. Peaks less than 0.1% of the TIC are not quantitated.

(d) Method Quantitation Limit (MQL) for agent is (50 ppm).

Appendix C 167

APPENDIX D

Gross Lesion Appearance (24-hr)

Project #: <u>G1555-</u>	38A	_				Date	2-210-6	<u> </u>
MREF Protocol #:	109		,		Study Di	rector: Carl Ol	son	
Day: <u>2</u>		Lesion	Read By:	<u>5</u>	Lesions !	Recorded By:	Dmm	
Lesion Sites	А	С	E	G	В	D	F	COMMENTS
Animal I.D. #								y
301	15/10	15/8	16/14	13/14	22/	19/33	13/20 R-3 E-3	readings taken in mm
	E432 R-2	R-2	E-3 R-3	R-2 E-2	R-3 E-2	R-2 E-2	R-3 €-3	
Mean Average								
					(OU) A	1 2-20-9	6 EMM	
All Measurements i N/A = Not applied N/II = Not require	ble	·s.	R = Eryther E = Edema 1 = Mild 2 = Moder 3 = Severe		•			
Stie A 1 Dul	ر دوار ۲۸ ز	_CHC	3					
Site B 50 L 1								
Site C 1000								
Site D 50.2								
Site E Car								
Site F 50.0								
Site G 1								
Reviewed By:				Date:	2/23	115		

Project #: <u>G1555-</u>	38A					Date:	2-20-	96
MREF Protocol #:	109				Study Dir	ector: <u>Carl Ois</u>	on	
Day: 2		Lesion	Read By: C	<u> </u>	_ Lesions R	ecorded By:	RMM	
Lesion Sites	Α	С	E	G	В	D	F	COMMENTS
Animal I.D. #								
305	108	12/2	9/6	14/3	19/20	23/	3125	readings takuin
	R-2 E-2	R-3 €-3	R-2 E-12	R-2 E-2	R-32	R-3 ⊆-3	6.65	
Mean Average								
II Measurements in IA = Not applicable /R = Not required in IA Not req	ole 1. 1.	<u> </u>						
ite B <u>50 nd 1</u>								
ice C <u>10nl 1</u> ice D <u>50nl 1</u>								
ine $E 10.0$			_					
to F_50.2								
ico G <u>lub</u>	town	47						
cviewed By:	176			Date: _	7/2	3/3/		

F Project #: <u>G1555</u>	5-38A					Date	<u> </u>	<u>96</u>
MREF Protocol #					Study Di	rector: Carl O	Ison	
Day: 2		Lesion	n Read By:	84	Lesions I	Recorded By: _	Omm	-
Lesion Sites	A	С	E	G	В	D	F	COMMENTS
Animal I.D. #								
306	15/8	7/8	78	16/7	16/3	12/4	112/0	realing taken
300	Ŗ- 2 Ε- 3	R-3 E-3	R-2 E-2	R-2 6-3	R-2 E3	R=32 E-2	R-1 E-3	
	-			-				
	 			 			 	
								·
				<u> </u>				
Mean Average								
		<u></u>			<u> </u>		OIF J-	22-96 RMM
All Measurements i N/A = Not lpg!jm N/R = Not required	مراط		R = Erythe E = Edema 1 = Mild 2 = Moder 3 = Severe	•				
Stie A 5.2.10	الله لما	-CHCI-						
Site B 1000 10								
Sice C 500 10	TO HD	in CHC	و آ					
Sice D 10, Q 10	Jo HD	inche	£					
Site E _ 5-210	7. HN	in CHC	ج لـــ					
Site F_10-210	090 tin	incHo	73					
sice o I me m		_						
Raviewed By:	17	<u> </u>		Date: _	2/23/	76		

	Project #: <u>G1555</u> -	-38A					Date:	<u> </u>	<u> </u>		
	MREF Protocol #:					Study Dir	ector: <u>Carl Ols</u>	on			
	Day: 2		Lesion	Read By:	BE	Lesions Recorded By:					
	Lesion Sites	A	С	E	G	В	D	F	COMMENTS		
	Animal I.D. #			1	1	1 -/					
	309	16/0	12/3	15/14	1410	79	9 9	7/0	taken in mm		
		R-3 E-3	R-1 E-2	R-3 E-3	R-1 €2	R-3 ⊭-3	R-22	E-3 E-3			
				_							
·-											
•											
	Mean Average										
l	All Measurements in N/A = Not applicable N/H = Not required	ole	· · · · · · · · · · · · · · · · · · ·	R = Eryther E = Edema 1 = Mild 2 = Moder: 3 = Severe				•			
\$	Stie A 10 22 1	107.H	DincH	<u>i</u> Cl 3							
	Site B <u>5-210</u>										
	Site C 1020 10										
	Site D <u>5~Q [</u>										
	ite <u> </u>							·			
	inc F 522 1										
	ice G <u>lus</u>										
R	civicwed By:	1 7	57/2-		Date: _	5/53/	<u> </u>				

Project #: G1555	5-38A					Date:	2-28	-96
MREF Protocol #	#: <u>109</u>				Study Dir	rector: <u>Carl Ols</u>	. កព	-
Day: 2		Lesion	Read By:	15	_ Lesions R	Recorded By: _	N-ALK-	-
Lesion Sites	А	С	E	G	В	D	F	COMMENTS
Animal I.D. #								
312	15-12	1910	14/2	20,5	9/2	311	12/3	Readings taken in mm
	R=-3	8-2 8-2	R-2 =:2	R-3 E-3	R-3 E-3	R-3.	C 3	
					;			
Mean Average								· ·
<u> </u>	<u> </u>				L	DWN 10	-23-96	D mn
All Measurements i N/A = Not applica N/R = Not require	bla		R = Erythen E = Edenia 1 = Mild 2 = Moders 3 = Severe					·
Stie A 10 ul 10								
Site B <u>5 ul 10</u>			_					
Site C 10.48 10								
Site D <u>5ul 10</u>		_						
Sice E <u>/Oul/O</u>								
Site F <u>5 H / 0</u>	To HU	in CHO	<u>~/</u> 3		-			
Site G <u>Lul A</u>	rest 1	HO						
Reviewed By:	178	115		Date:	2/25/8	6		

	Project #: <u>G1555</u> -	38A					Date:	2-25-9	<u>.</u>
	MREF Protocol #:	109				Study Dir	ector: <u>Carl Ols</u>	sonno	
	Day: 2		Lesion	Read By:	70	_ Lesions R	ecorded By: _	North	
	Lesion Sites	А	С	E	G	В	D	F	COMMENTS
	Animal I.D. #		,						
	316	13-9	15/15	314	3314	11/9	15/9	14/14	Riadengo taken mm
		R-2	8-3	R-3 E-2	R.3	R-2 =3	R-3 £ 3	R-3 E-2	
									·
			<u> </u>						
						<u> </u>			
	Mean Average								
	All Measurements in N/A = Not applicat		s	R = Erythem E = Edema	na	•			
	N/R = Not required			1 = Mild 2 = Modera	.te				
ç	Stie A 10 sel	10% 1	4NinC	3 = Severe					
	Site B Jul /								
	Site C 16 11 /1								
	Site D_Ful /C								
	ite <u> </u>						·		
	ite F <u>-52Cl//C</u> ite G <u>-/2Cl/</u>			<u>_/</u> _3					
						_ -			
R	eviewed By:	()	C15-	<u>. </u>	Date: _	2/23/	<u>) [</u>		

Appendix D

174

	•		ON SIZE	E DETFI	RMINA	TION SI	HEET			
Project #: G15	6 ³⁸ / 55 -200	 					Date:	3-6	-96	
MREF Protocol #: 109 Study Director: Carl Olson										
Day: Lesion Read By: Lesions Recorded By:										
Lesion Sites	А	С	E	G	В	D	F	н	COMMENTS]
Animal I.D. #			<u>.</u>		(Ð				1
313	1013	10 g	13/14	14/16	NANIA	NANIA	NAJA	NANA	reading to	
	R- 2	R-2 E-3	R-3 E-3	R-3 E-3		K-0 1 E-1		R-0 ' E-1		
					2-0 E- 1	K3) Tart				
										•
Ll									W.U 3-6-96	Aire
All measurements in N/A = Not applicab N/R = Not required	le		E = 1 $1 = N$ $2 = N$ $3 = S$	Erythema Edema Mild Moderate Severe	0 = @:	Ac Notapa AC Non On	carcut 1A is this it t	egu v formate ct	ivalent to m throw indy 10-24	1. 00 gh 1-96 Oas
Site A 5 11 10%	HD	in CH	2/3				_		. <i>U</i>	
Site B 20ul ne	trali		eletion	\sim			•			

Site A <u>5ul 10% HDin CHC13</u>

Site B <u>20ul neutralizing</u> solution

Site C <u>5ul 10% Hamin CHC13</u>

Site D <u>20ul neutralizing</u> solution

Site E <u>5ul 10% Lin CHC13</u>

Site F <u>20ul neutralizing</u> solution

Site G <u>1ul neutralizing</u> solution

Site H <u>20ul neutralizing</u> solution

Form No. MREF-LESION.SIZ-07

OENTHUELLOND 3-6-176 SALH
BEE 13-6-46 JARH

Appendix D

Revenuel by CT Dian

	•		ON SIZE	E DETE	RMINAT	tion sf	HEET			
Project #: G15	0384 55- 9001						Date:	3-6	-96	
MREF Protocol	#: 109)		_ Stud	ly Direct	tor: <u>Car</u>	l Olson			
Day: 2	L	esion Re	ad By:	<u>(† 5</u>	L	esions R	ecorde:	d By: 🗻	JncH	
Lesion Sites	A	С	E	G	В	D	F	Н	COMMENTS	
Animal I.D. #			<u>, </u>			2)	·			
315	150	12/13	11 9	18/12	NIANIA	NIA	NIPLIA	NIANIA	reading the service	
	P-1 F-1	L-3 E-3	R-2 E-3	と-ス E-ス	E-1	R-01 E-1	2-0 E-1	2-0' E-i		
										•
All measurements in N/A = Not applicat N/R = Not required		ers		Erythema Edema Mild		0	= 47 63 AC 1	tappa NAIST	rent OWN 3- equivalentha Growtha Hidy 10-24	6-96 On It to ?
1111C = 1100 104anaa			2 = 3	Moderate Severe			₹		Didy 10-24	-96 DA
Site A <u>Dul 10 %</u>	3 HD).	mCH(213							
Site A <u>. T. ul. 10 %</u> Site B <u>20 ul. M2</u> 1	utral	incre s	colutio	7						
ein a 5 11 0 10%										

Site B20ul neutralizing solution

Site C5ul 10% Lin CHC13

Site D20ul Mentralizing solution

Site E5ul 1070 HD enCHC13

Site F20ul neutralizing solution

Site G1ul neutralizing solution

Site H20ul mentralizing solution

Form No. MREF-LESION.SIZ-07

Reviewed by C7 Bl s-

LESION SIZE DETERMINATION SHEET										
Project #: G15	Ф 33.4 55- 900						Date:	3-6	6-96	
MREF Protoco	1#: <u>10</u>	9		Stud	dy Direc	tor: <u>Ca</u>	rl Olsor	1		
Day: Lesion Read By: Lesions Recorded By:										
Lesion Sites	A	С	E	G	В	D	F	Н	COMMENTS	
Animal I.D. #	Animal I.D. #									
317	9/5	8 8	13/13	15/10	20	90	20	90	Taken MIL	
	2-3 E-3	12-3 E-3	R-2 E-2	1-3 E-2	R-0	R-0 E-1	R=0 =-1	R-0 E-1		
										,
All measurements in N/A = Not applicable N/R = Not required		ers	E = 1 1 = 1 2 = 1	Erythema Edema Mild Moderate Severe	0=	Ylot a	ppare	F & 27	UN 3-6-96 B. E 3-6-96 B.	MR Ma
Site A 548/07	Lim	CHC/3								
Site B 20ul ne	utrali	ama a	elution							
Site C <u>5ul 10%</u>										
Site D <u>20.ulnc</u> ı	itial:	yers s	election	·						
Site E <u>5 ul 10%</u> 1		-								
Site F Quul Me	utral	izing.	soluti	% →~						
Site G <u>ful ne</u>										
Site H <u>20 ul m</u>	intra	lizing.	sodiete	سيس						

Appendix D

Form No. MREF-LESION.SIZ-07

177

Reviewed by 67 Clar-717196

'D -1# C15	0354	:						2	/ 0.
Project #: G15)) -700						Date:	<u> </u>	2-96
MREF Protocol #: 109 Study Director: Carl Olson									
Day:	I	esion Re	ead By:	<u> </u>	L	esions R	Recorde	i Ву: _	Jm _H
Lesion Sites	A	С	E	G	В	D	F	H	COMMENTS
Animal I.D. #									
324	9/12	10/11	13/14		00	200	26	20	telen mm
	E-3 E-3	2-2 E-2	R-3 E-3	E-3	R-0 E-0	P-0 E-1	R-0 E-1	R-0 E-1	
				}					
1.									
All measurements in N/A = Not applicab N/R = Not required		rs	E = 1 1 = 1 2 = 1	Erythema Edema Mild Moderate Severe	Ð =	Not ap	parent		UN 3-6-96 DAN
Site A 5 ul 1070	HDi	wCHC.	13						
Site B 20 1 Mes	trali	sing so	olution	-					
Site C 5 2 107, 1									
Site D 20 ul neu	itrali	sing Di	olution	•					
Site E <u>5ul 107</u> 21		<i>O</i>							
Site F. 20 al Mer.			lution	_					
Site G Lel nea	t HI) 2							
20.0			1-1-	•					

Revenued by CT DRS-

Form No. MREF-LESION.SIZ-07

31

Appendix D

Project #: G15	-35.1 55-9 001	0	51. 012.		.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	. 1011 01	Date:	3-6	6-96	
MREF Protocol				_ Stud	ly Direct	or: <u>Car</u>	l Olson			
Day: _2	L	esion Re	ad By:	Car	L	esions R	ecordeo	i By:	IncH	
Lesion Sites	A	С	E	G	В	D	F	н	COMMENTS	
Animal I.D. #										
3//	13 11 2-2 E-2	14 9 R-3 E-3	13-9 R-3 E-2	15/10 R-2 E-2	00 R-0 E-1	00 R-0 E-1	0/0 1-1	0/C R-0 E-1	reodings teamin mm	
								-	•	
				: بیر						
										0.00
All measurements in N/A = Not applicab	ole	ะเร	E = 1 = 2 = 2	Erythema Edema Mild Moderate Severe	0=	: Leta	rppai	ent a	3-6-96 לגעו 1 מינט 3-6-96	oma
Site A 5 w 10%	HNii	<u>.C.HCl3</u>	1							
Site A <u>5 ul 10%</u> Site B <u>20 ul Ne</u>	itrali	ging si	lution							
Site C <u>5 ul 1070</u>										
Site D. 20ul ne		. ,	colution	i.L						
Site E <u>5 4 6 10 76 1</u>	H Din	<u>CHC/3</u>	,							
Site F 20ul 126				ion						
Site Goul ne										
Site H <u>ZOUD</u>	eutra	lizing i	من من مناها	r						
Form No. MREF-LESIO	N.SIZ-07									<i>C</i> 3

Residency 60 CT 6085-

Project #: G15	55- 38A	7				Date:	<u>.3-</u>	14-96	
MREF Protocol	#: 109	Ph	<u>.ur. 3</u>	_ Stud	dy Direc	tor: <u>Ca</u> r	i Olson		
Day:2	L	esion R	ead By:	Æ_	L	esions R	Recorde	d By: <u>LOM</u> ,	<u>ku</u>
Lesion Sites	А	С	E	G	В	D	F	COMMENTS	
Animal I.D. #	0								
310	#135		12/0	910	1722	11/17	15/6	taken in ma	
	R-1 E-2	R-1 E-2	R-3 E-3	E-3	R-1 E-2	R-1 E-2	R-05-Q	·	
					;				
					•				
All measurements in N/A = Not applicab N/R = Not required		ers	R = E = 1 1 = 1 2 = 1	Erythema Edema		OAC Sust Sustre Leve	the le deter vew p. fore	mined at 0 and lan designation	dingo Were livelod id was of as 0.5. 96 pum
Site A <u>5 ul 107</u>	HDin	_CHC	13	•	(BEE 1	0-16-96	ELM	
Site B 25 ul Re									
Site C <u>Jul 107</u> 2	HNin	_CHC	3						
Site D <u>25ulBl</u>			_						
Site E <u>5 ul 10%</u>	Line	CHC13							
Site F <u>25ulCh</u>				1~,—				·	
Site G <u>/ul_ns.</u>	et H	<u> </u>							
Form No. MREF-LESION Appendix D	i.SIZ-07			180)		ب کون دور	سیل کمس ^د ۱۱۶۱۶ه	T DESL

						_	_		
Project #: G15		_						14-96	
MREF Protocol	#: 109	Pha	<u>u 3</u>	Stuc	ly Direct	or: <u>Car</u>	l Olson		
Day:	I	esion Re	ead By:	OB	L	esions R	ecorde	i By: <u>RM</u>	<u>~_</u>
Lesion Sites	А	С	E	G	В	D	F	COMMENTS	
Animal I.D. #									
491	12-9	11/15	3 10	13-19	7/2	15/1	9/26	reactings	
	R-3 C-3	R-3 G-2	R-1	R -1 ≤-3	R 05-64	R-1 E-2	P.31 5.3.		
									
·									
					,				
				j					. ·
All measurements in in N/A = Not applicable N/R = Not required Site A Sul/07 Site B 25 ul Cha Site C Sul/07 Site E Sul/07 ite F 25 ul Bl ite G / ul Maa	Limi vicoal HDim d Wa HVin	CHC13 Wasta CHC13 Latest CHC13	R = 1 E = 1 1 = N 2 = N 3 = S	violerate levere ite Ar tim pu at on luy nu	aday seid she can a care a car	1 app ick a ick a ic	2-96 A 2-96 A 2-96 M 2-14-96 ma e 2-14-9 5-14-9	the lesion the lesion para Col notected saletran Col nimels of le Drin adinos levelis was the	were beet between

Form No. MREF-LESICH.SIZ-07

Appendix D

181

Revenued by CT OPSIL

Project #: G15	55- 38A					Date:	3-19	1-96		
MREF Protocol	#: 109	Piña	<u> </u>	_ Stud	y Direct	or: <u>Car</u>	l Olson			
Day:2	L	esion Re	ad By:	<u> 16</u>	L	esions R	ecorded	I Ву: <u>Дл</u> и	<u>ıc</u>	
Lesion Sites	A	С	E	G	В	D	F	COMMENTS		
Animal I.D. #			·		,	y				
493	9/7	11/7	19 4	14/15	150	19 19	15 20	. Readings taken in MM		
	K-3	6-1 6-2	R-3 E-3	R-3 E-3	R-1 E-i	R·1 €·2	R 0.50 2-05			
								,, - ,- ,- ,- ,- ,- ,- ,- ,- ,- ,- ,- ,-		
					<i>:</i>				••	
									í	
All measurements in N/A = Not applicab N/R = Not required		ers		Erythema Edema Mild	OAC d	the le eterm ,0 que	sioni ined LI an	st livels dwart	were best between herefore 7-14/96 DM:	٠,
	4			Moderate Severe	0	leotar	كستكث	د دان بنده	, , ,	
Site A <u>5 LU 109</u>										
Site B <u>25ul <i>R</i> :</u>										
Site C. <u>5 ul 107</u>	_									
Site D <u>25ulBl</u>	uc W	acteut	مصمعة							
Site E 5 ul 1076	Lin	CHCI	3							

Form No. MREF-LESION.SIZ-07

Site G I w neat HD

Site F 25 ul Charcoal Wastertream

Appendix D

Reviewed by CT Elsa 3/18/50

Project #: G1555-38A Date: 3-14-96										
MREF Protocol	#: <u>10</u>	P	hase. 3	Stud	iy Direc	tor: <u>Ca</u>	ri Olson			
Day:2	I	esion Re	ead By:	16	L	esions R	Recorde	d By:C_M	<u>u</u>	
Lesion Sites	A	С	E	G	В	D	F	COMMENTS	7	
Animal I.D. #									1	
498	79	12/1	910	17/5	1733	1100	17/9	Asadingo takinin mm]	
	R-3 E-1	R-3 E13	2.3 :3	R-2 C-3	R-3 E-i	662 E62	6-2			
			7.1							
								: 		
·										
										
									·	
All measurements in N/A = Not applicab		ers	E = 1 1 = 1 2 = 1	Erythema Edema	(diters	nine	eat level	between therefore 3-14-46 PM	•
Site A <u>Sul 107</u> 2	HNin	<u>CHCI</u>	3							
Site B <u>25ul Blu</u>	بالكم	ctestre	on							
Site C <u>5 ul 10%</u>	Lin	CHC13								
Site D <u>25ul Cha</u>	rcoal	Wasti	stress							
Site <u>E. 5. ul. 10%</u>	HDin	CHC1	3							
Site F25ul Red										

Form No. MREF-LESION.SIZ-07

Site G Lul Mest HD

Appendix D

Revenied by CT Dism 3/15/94

		FF216	ON 312.	E DETE	RMINA	LION SI	HEET		
Project #: G15	<u>55- 38</u> 4	. 4				Date:	3-6	2-96	
MREF Protocol	#: <u>10</u> 5	2 Pho	1023	Stud	dy Direc	tor: <u>Car</u>	l Olson		
Day:2	I	esion Re	ead By:	_A3	I	esions R	tecorded	i Ву: <u>ДОм</u>	m
Lesion Sites	A	С	E	G	В	D	F	COMMENTS	
Animal I.D. #									7
499	78	1010	11	12/6	13/5	15/6	9/6		7
	R-2 E-2	R-2 E-3	R-3 E-1	Q-3 ≤-2	Q-1	R-0.5	R-1 E-2]
		U=40	u=6	u= 4					1
									7
					·				
	•		·						
All measurements in	millimete	rs ()=mal R =	o planin Erythema	TOAC	. The	lesio	n reader	go were best velo between
N/A = Not applicab N/R = Not required	le		E = 1 $1 = 1$	Edema Mild		معت	ermen Ond	did at li	a thurfore
·				Moderate Severe		معك	ugra	Ted as on	e therefore 5. 3-2246 Rr
	*	· . ~ 4			(T) 1				•
Site A <u>5 ul 107</u>					(J) H	عامر ال عالما	ren	et pewi	ose sites ously records 96 DMM
Site B 25 (_			
Site C 5 11/0°					U =		iction	u noted :	۵0:
Site D_25 ul C-	haice	ial Wa	stiler	·		5	- mio	liem	
Site E <u>5 ul 10°</u>	7 HD	rin Cl	4C/3			6	مىمد	ge .	
Site F 25 ml K	ed h)asti	dream	<i>.</i> _					
Site G/ul Me.	at H	<u>'D</u>							

Form No. MREF-LESION.SIZ-07 Appendix D

Reviewed by CT Else 3/28/96

		LESIG	ON 2121	E DE CO	'Q VIINA	110N 21	HEET		
Project #: G15	55- 38A	7				Date:	3-2	2-96	
MREF Protocol	#: <u>109</u>	Dh.	ase3	Stud	iy Direc	tor: <u>Ca</u>	rl Olson		
Day:2	L	esion Re	ead By:	<u>Æ</u>	I	esions F	Recorded	i Ву: <u>Ют</u>	<u>-m</u>
Lesion Sites	A	С	E	G	В	. D	F	COMMENTS	
Animal I.D. #									
494	14/3	10/3	10/1	17/5	00	21/19	1320		1.
	R-3 E-3	R-2 E-2	R-2 E-2	R-3 €-3	R-0	R-1	R-2		
			U=4	4-4					

·									
					,				
All measurements in N/A = Not applicab N/R = Not required	le		E = 1 = 1	Erythema Edema Mild Moderate Severe		O=M OAC - Mot U=	ot ap ulcer pres ulcu	ation of a viously 3.	lose setes w recorded 28-96 orim reter an);
Site A <u>5ul 10</u>	70 Li	<u>nCHC</u>	2/3				3	4 = smal 5 = medie	in
Site B 25 ul Ch	arcoc	Dwa	steatr	ion		•	4	:= large	,
Site C <u>5ul 10</u>	70 H C	rin CH	K13						
Site D <u>25 ul</u> R				,					
Site E <u>5 1 U 10°7</u>	<u>, HN.</u>	<u>in CH</u>	10/3						
Site F <u>25ul Bl</u>	ue h	astes	trean	U					
Site G <u>/ul</u>	eat b	10_							

Form No. MREF-LESION.SIZ-07 Appendix D

185

Revocat by CT Dem

Project #: G15	55- 38A	7			Date: 3-22-96					
MREF Protocol	#: 109	Pho	<u>er 3</u>	_ Stud	ly Direc	tor: <u>Car</u>	l Olson			
Day:2	L	esion Re	ead By:	<u></u>	L	esions R.	ecordeo	i Ву: <u>Ю</u> пип	<u>u</u>	
Lesion Sites	А	С	E	G	В	D	F	COMMENTS		
Animal I.D. #										
496	12/5	109	13/6	17/6	1921	1919	00			
	R-3 E-3	R-3	R-3 5-3	K-3	R-1 E-2	R-3 E-2	R-0		*	
	u=6	سدد	U=6	W=4						
·										
				-					. •	
					,				·	
									_	
All measurements in N/A = Not applicab		ers	E = 1 = 2 = 3	Erythema Edema Mild Moderate Severe					t, of dose viously 896 RTMM	
Site A <u>5 ul</u> 107	6 HD	in CHC	! /3					4=2	mall nedium	
Site B 25 ul R	ed Wo	estest	eim			•			neduum arge	
Site C <u>5 ul 10</u> 6	70 HL	<u> Iin CH</u>	C/3					6-2	ary	
Site D <u>25 ul</u> <u> </u>	<u>Me W</u>	actest	Team							
Site E <u>52l 107</u>	70Lin	<u>~CHC</u>	/3							
Site F <u>25 ul (-</u>				bream						
Site G <u>ILL</u> M										

Form No. MREF-LESION.SIZ-07
Appendix D

186

Received by Ci Olse-3125192

•						•			
Project #: G15	55- 38A	۷.				Date:	3-2	2-96	
MREF Protocol	#: <u>10</u> 9	2 Ph	<u>au 3</u>	_ Stud	iy Direc	tor: <u>Ca</u>	rl Olsor	L	
Day:2	L	esion Re	ead By:	<u> 9</u> B	I	esions R	Recorde	d Ву: <u>Д</u> м	<u>M</u>
Lesion Sites	A	С	E	G	В	D	F	COMMENTS	
Animal I.D. #									7
497	8 8	11/10	10/12	1521	2120	1815	14/18		
	હું. જુ. જુ.	R-3 E-3	R-3 =-3	R-3 G-3	R-2 G-2	R.O.5 E	R-2		
	u=60		U=68	لدعدج]
								·	
					<u> </u>				
					· ;				
					· ·				
									OIE 3-22-96 BM
All measurements in N/A = Not applicab N/R = Not required		rs	1 = 1 $2 = 1$	Ettema		Lot H	tive dieef	n 0.0 and one design 3-22-96	ngo were at levels - I and was mated as
Site A <u>5 wl /07</u>					(3)	AC L	ilcera vece .	ation of co	lose actes viously record
Site B <u>25 sel B.</u>	ene w	<u>acies</u> i	مسمدوع			•	9-0-0	5-76 NG 12C/N	
Site C <u>5111 107</u> 5	Lin	<u>,CHC</u> ,	13		l			tion not	iclav:
Site D <u>25 ul Ch</u> a	nconl	West	estria.				med		
Site E <u>5 nd</u> 1070	HDI	m CH	(C/3				lar		
Site F 25 ul Re	ed We	estesta	edur						
Site G / ul m	eat 1	4D							

Form No. MREF-LESION.SIZ-07
Appendix D

Revenued by C. T. 6285-

Project #: G15	Project #: G1555-38A Date: 6-21-96											
MREF Protocol	#: <u>10</u> 9	Pho	<u>E</u>	Stud	iy Direc	tor: <u>Car</u>	-l Olson					
Day: 2	L	esion Re	ad By:	612	I	esions F	Recorde	d By:	D.pcm			
Lesion Sites	А	С	E	G	В	D	F	Н	COMMENTS			
Animal I.D. #				_		,						
346	28/2	12 8	10 g	11/14	00 R-0	16 21 R-1	00	19/20				
	R-3 E-3	E-2	E-1	E-1	E-0	E-1	E-0	€-1				
						ļ						
					,							
All measurements in N/A = Not applicab N/R = Not required		ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe			= Me	t opp	prent			
Site A <u>5 1 1070 1</u>	Linc	4013										
Site B 25-2-	te B 25 ml mattation											
Site C 5-4 1070	~ <i>I</i> #	<u> </u>	3									
Site D 25 2 62	ea mo	- teate										

Site H 25 al Some mosterine

Form No. MREF-LESION.SIZ-07

Site = 25 a motost

Site G Indust HD

Site E 5 2 10 10 10 10 in CHC13

Appendix D

Reviewed by UT ODE 6/24/96

							_		•	
Project #: G15	55- 38A	7				Date:	<u>e-</u> .	21-96		
MREF Protocol	#: 109	Phs	<u>3</u>	Stud	dy Direc	tor: <u>Ca</u>	rl Olson			
Day:	L	esion Re	ead By:	_B11	I	esions F	Recorded	i By:	Emm	
Lesion Sites	А	С	Е	G	В	D	F	Н	COMMENTS	1
Animal I.D. #	}					_			1	
341	11/2	11/8	14/3	159	2413	0/0	11/12	0/0		
	R-2 E-2	R-2 E-1	R-3 E-3	R-3	R-1 E-1	Q-0	2-3 E-2	E-0		
	u=50	u=4 0	426 D				4-60			
					ļ	ļ				
All measurements in N/A = Not applical N/R = Not required	ole	ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe	. (C)	AC III Ne Dwo	lceraticers	tion of	pporent of dose sit restivusly 34-96 Don	teci uu
Site A 5-2107	· HD	-CHC	£				a	1 =	<u>+</u> · d = 5	7.
Site B 25.0 60	صدعب	tate			بر ز	1 = M.	מימשל איי	ttor .	neted as	
Site C 5-210%	HWin	<u>-CHC</u>	3					3	5= midues	
Site D 25	7	tate						(6=large	
Site E 5-2 (07)	<u>o Lin</u>	CHCL.								
Site F 25.0 ba	ئىوسى_	<u>artarte</u>								
Site G 1 22 00	计大	7								
Site H 25-2-	<u> </u>	سلتسك								
Form No. MREF-LESIC	N.SIZ-07						Rei	بمعدرسيا	by	

Appendix D

iù.

189

Covering by

Project #: G1555-38A Date: 6-21-96										
MREF Protocol										
Day: 2	I	esion Re	ad By:	B/	<u></u>	esions F	Recorde	d By: <u>→</u>	Omm	
Lesion Sites	Α	С	E	G	В	D	F	н	COMMENTS	
Animal I.D. #										
339	109	18 13	12/12	9/10 R=3	0/0 R-0	19/14 R=2	00	28/13 R-2		
	E-1		E-3		E-c	<u>=-2</u>	E-D	R-2 E2		
		4-40	U=40				<u> </u>			
					-					
					3					
					-					
All measurements in N/A = Not applicat N/R = Not required	ble	ers	E = 1 = 2 = 1	Erythema Edema Mild Moderate Severe	©	AC 2 2	باروی پیورو پیورو	ation net ded	egarent of does s frevious 6.2:-96 d	
Site A 5-010	1H de	مندحه	tcl3			<i>il</i> =	· who		مسكفاءر س	-
Site B 25.2 ^									- small viedeur	
Site C 5-110°		_							Large	
Site D 25~2 b	مسع	tota	٠						,	
Site E 5-0-10	The HD	:. <u>C</u> #	تلء							
Site F_ 25_0 ~	7-	estreta								•
Site G Ludries	开土									
Site H_ 25.2 b	عده	motocu	Trans	-				Reuren	sid by	ŀ
Form No. MREE-LESIC	N.SIZ-07							6 6	illian ul	24/36

Form No. MREF-LESION.SIZ-07
Appendix D

190

Project #: G15	55- 38 <i>6</i>	7				Date:	6-	<u>21-96</u>	<u>-</u>
MREF Protoco	1#: <u>10</u> 9	Ph-	<u>423</u>	_ Stud	dy Direc	tor: <u>Ca</u>	rl Olson		
Day:	I	esion Re	ead By:	BH	<u> </u>	esions F	Recorde	d By: _	QNin
Lesion Sites	A	С	Е	G	В	D	F	н	COMMENTS
Animal I.D. #				· · · · · ·		.			
342	22 14 12-3 15-3	R2 E-2	9/10 R-2 E-2	15 g E-2 E-2	23 15 2-2 =-1	0 R-0	11 R-2 E-2	00 R-00	
				-					
			,	<u> </u>					
All measurements in N/A = Not applica N/R = Not required	ble	ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe			0 = 1	not c	epparent
Site A 5-210°									
Site B Salk									
Site C <u>5.2107r</u>									
Site D <u>25.l.v.e</u> Site E <u>5.l.l.O</u>									
Site F 25, 26		•	_						
Site G \								Rin un	سند لم
Site H 25.2									108m 6/24/
Form No. MREE-LEST	N 517-07								

Appendix D

Project #: G15		_		·				27-96	2	
MREF Protocol	#: 109	Pi	<u> 2263</u>	_ Stud	y Direc	tor: <u>Car</u>	l Olson			
Day:2	L	esion Re	ad By:	<u>El</u>	<u>+</u> L	esions F	Recorde	d By:	onini.	_
Lesion Sites	А	С	E	G	В	D	F	н	COMMENTS	
Animal I.D. #						z		4 6		
340	12/1	8 8	11/14	11/14	19/13	19/14	122/6	00		1
	R-2 E-2	R-2 E-2	R-3	是五	R-2 E-1	R-1 E-1	R-1 E1	R: 0		-
	U=4m			17:60						-
							<u> </u>			
										-
										•
					:					
All measurements in N/A = Not applicat N/R = Not required	ole	ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe	(DACJ			ot appa or dos	
Site A 540 107	O HD.	in C	40/3			W	، سىد(mits	127-9	y record
Site B 25_11 _1	Sluce	wast	estrec	27			•			
Site C 524/60					į	بار د د	Llcer 11-	ation	noted.	، نەد
Site D 25 1 L 1	eddi	racte	2012	ب <i>د</i>				med		
Site E-5 xcl /6							6 -	large	シ	
Site F 25 ul. &	<u> Luci li</u>	Jaste	atua	жU						
Site G / Lel m	int	HD_								•
Site H 25 ul 1	redui	acte.	Trea	yu-		Rue	سسولإ	by C	7 860-	L [25 SL

Form No. MREF-LESION.SIZ-07

Appendix D

Project #: <u>G15</u>	55- 38A					Date:	L, - a	27-96	<u>:</u> _	
MREF Protocol		\sim	<u>see 3</u>	Stud	y Direct	or: <u>Car</u>	l Olson	L		
Day:	L	esion Re	ead By:	BH	L	esions R	ecorde	d By: 💆	ampl	
Lesion Sites	А	С	E	G	В	D	F	н	COMMENTS	
Animal I.D. #				·			·			
345	9/13	914	11/10	14 15	00	1915	21/5	2016		
	K-3	R-3 E-3	R-2 E-3	R-2 E-3	R-0 E-0	R-2 E-1	R-1	R-2 E-1		
	14-6 E	لاءل	U=6	6-6 ()				U=4		
·										
							·			
				٠.						
										
All measurements in N/A = Not applica N/R = Not required	ble	ers	E = 1 = 2 = 1	Erythema Edema Mild Moderate Severe	(I) AC .	ulce not p		not appare	
Site A <u>5ull 10</u>	70Li	~ ette	13							0
Site B 25 ul x	<u>ed li</u>	laste	Itea n	_~	ب	در = رز	يلأوف	ration	v_notecla	بيم
Site C. 5 ul /0	70 HD	in C-t	10.13				4=-	ilmal medi	ث	
Site D.25ul	lun i	<u>vost</u>	The as	· · ·				large		
Site E. 5 ul 10								•		
Site F 25 ul 1			trea	m						
Site G /ul m					Ω		1	(D. 11-5	ls:
Site H 25 rd &	-lus-1	<u>want</u>	eatria	m:	Ke	ىشىرى ى قالىد	در لصس	()	Des 6/25	[1>

Form No. MREF-LESION.512-07

Appendix D

1

								.a.		
Project #: G15		_						27-96	<u>:</u>	
MREF Protoco	l#: <u>10</u> 9	ρ	have	Stud	ly Direc	tor: <u>Ca</u>	ri Oisor	L.,		
Day:	L	esion R	ead By:	_ B13	L	esions F	Recorde	d By:	<u>Chini</u>	
Lesion Sites	A	С	E	G	В	D	F	н	COMMENTS	
Animal I.D. #										1
351	13/5	159	10/15	15/14	16/19	15/2	2/12	00	1	1
	R-2 E-2	R-3 G-3	R-3 E-2	K-3 E-2	R-1 E-2	R-1 E-1	R-1	R-0 I0		
		12-4 ₀	11-Z	4-5 ₀						
·										
										•
All measurements in N/A = Not applicable N/R = Not required	ole		E = 1 1 = 1 2 = 1 3 = 5	Erythema Edema Mild Mioderate Severe	(DAC.	liêc Sere		net appa n Holos psolvious 6-27-961	
Site A <u>Jul 107</u>	OHN	in CH(43				/UC	rded	627-961	G perc
Site B <u>25ulh</u>	luz l	saste	stream	~		U =	ulc	erati	in note	dae:
Site C <u>5 ul 10°</u>	<u>るんこ</u>	<u>nch</u>	213					= xin		
Site D. 25 ul ru	ed w	aste	itreas	~				lan	dium ge	
Site E. 5 ul 10%	TO HD	<u>in C</u> t	1013						•	
Site F <u>25 1 L. E.</u>	luit	Daste	alria	yn.						
Site G <u>/ul ma</u>	est.	40								
Site H_25_ul_1			etre à	yn		2	,	h. 0	T Des	6[25[5]

Form No. MREF-LESION.SIZ-07
Appendix D

194

Project #: G15	55- 38A	7				Date:	6.6	77-96	<u>:</u>
MREF Protocol	#: <u>109</u>	\mathcal{L}_{-1}	lase.3	Stud	ly Direc	tor: <u>Car</u>	l Olson		
Day:	L	esion Re	ad By:	BH	<u> </u>	esions R	ecorde:	d By: 🔥	912-pm
Lesion Sites	А	С	E	G	В	D	F	н	COMMENTS
Animal I.D. #									
3.52	119	12-9	- 1 V	13/15	00	16/15	15/2	15/9	
	R-2 ≂-3				R-0 E-0		R-1 E-1	R-2 E-i	
	ند- ع	11-4 F		U-6					
									·
								C= m	t apparent
All measurements in N/A = Not applicab		rs	E = I $1 = N$	Juderate	Œ 1º			6 Emi	

Site A Jul 10% HD in CHC/3 Site B25ul rid Wastatrian Site C 5ul 1070 HNin CHC13 Site D 25 ul blue Wastestrom

Site E 5 4 10 70 Lin CHC 13

Site F 25 ul red Wastestream

Site G/ul mest HD

Sice H. 25 ul blie Wastestream

Form No. MREF-LESION.SIZ-07 Appendix D

QAC alceration of dose sites were not perviously recode 6-27-98 Amm

U- ulceration noted as:

4- small

5- medium

6- lange

Reviewed by CT Dem 6/28/86

Day: 2	L	esion R	ead By:	013		Lesions R	ecorde:	d By: .	DININ_
Lesion Sites	A	С	E	G	В	D	F	H	COMMENTS
Animal I.D. #			7 	1 =		4.6	(., -)	T	
383	15/20	11/18	1918	0	6	10	16/4	0	
	K-3	5-7 5-7	R-2 E-2		G-7	R-,53	K-1 ⊈-,		
					_				
· .									
									-
					<u> </u>				
				1					
I/A = Not applicable I/R = Not required ite A 10 ul 10 ul ite B / O ul bl	7. L.		1 = 2 = 3 = C/3	Edema Mild Modera Severe	te				
ite C 10ul 10°									
ite D 10112 Che				treas	v				
ite E/Oul 107									
ice Flour Nic				100				D	· ~ Mh ~ ~ ~
te G_	~~~	مسمت		,,,				المحا	مر ورار الا
to I!									
									

Project #: G15		_						: <u>8-</u>		96	_	
MREF Protocol	#: 10	2 Ph	0023	_	Study	Direc	tor: <u>Ca</u>	rl Olsor				
Day:2	I	esion R	ead By:	R)	1	Lesions 1	Recorde	і Ву	: <u> </u>	<u>Omm</u>	
Lesion Sites	A	С	E	G		В	D	F]	 H	COMMENTS	
Animal I.D. #		·										
385	13/5	13/8	12/16	0		2/12	15/7	1/4	0			
·	R-1 E-2	R-3 E-2	R-1			-2 -2	R-1	R-,50				
										1-1		
				\dashv						1+		
					+					$\frac{1}{1}$		
										H	i	
				+	+					\vdash		
N/A = Not applicable N/R = Not required	•		E = I $1 = N$ $2 = N$ $3 = S$	fild forlera								
Site A1011107	HN	мCH(13					,				
Site Blow Ald	اللاها	state	an									
Site C/Oul 100	6 Li	-CHO	213									
Site DIOul Bl	-			_								
Site E 10.48 10°7									D		16 (7	7
Site F10 we Che				سمم					الع	مکا لہ	welly C	71+1
Site C												
(APPENDIX D) FORM NO. MREF-LESION. CAC These	.\$12:07 _ OLL	 اس سا	مر عاب	19 شي	7 المعالم	ed:	or F	. d	ec	8-	13-96 Rm	~ <i>}</i> ;^
@IF 8-14-96 (3)AC Lice less C.O and	DM,	n. readi	ا تهرسه شهر	1	ربوء المحاد	یک ط مسا	itim mati	uned was	at o c	اعثاد . سح.	vela bletw 8-14-96 Pi	سور المرام

Project #: <u>G1555</u>	5- 38A						Date:	_8-	14-	-91	2
MREF Protocol #	t: 109	Pa	ase?	3	Stud	dy Direc	tor: <u>Ca</u>	rl Olson			
Day: 2	L	esion R	ead By:	B	<u>+</u>		esions R	Recorde	d By	: _/	<u> Emn</u>
Lesion Sites	A	С	E		G	В	D	F	1	H	COMMENTS
Animal I.D. #				,							
400	77	15/5	1521	0		5,9	11-14	13-14	0		
i i	ارا اور الله اور	R-3 F-3	R-3 C-3			E-02	R-1 G-0	R-1 E-1			
										_	
		· · · · · · · · · · · · · · · · · · ·								-	
`									·		
					:						
Site A Dul 1076 Site B 10 ul Cha	rco. HN.	e wa in CF	3 = 1 HCl3 stestro ICl3	ean	e						
Site D/Oul rec											
Site E 10 ul 10%											
Site F/Och blu	<u>دنار و</u>	arle	"Learn	-				0			. , , , , ,
Site G_								K	عاث (سند	2 by 0.7.0
- Site II		<u> </u>									
(Appendix Form No. MREF-LESION.	D) s12-07			1	.98			(3) .	,		0 12-96 RA
(Appendix FORM NO. MREF-LESION. AC The lead O.O and WICE STEEL S	ette	i lili Neadi	nger i	رن-ا رموءا سين	- ك-	enice Bent One	deter	ای این (سی) انتخاری که ۱۳۶۰ د و در در در	el ! a. ! o	سند کرکے دور	60.76 8-14-96
0.0 and	.1 a	سط که -	jau 7	<u>ا</u>	ا		Lung	،سر دد	- 0	-	

	Project #: G15	<u>55- 38</u> A	٠ .				Date:	8-	14-	96	
	MREF Protocol	#: <u>10</u> 9	Pz	رى د تى	3 Si	udy Direct	or: <u>Car</u>	l Olson			
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APPENDIX E

Dosage Site Code and Histopathology

Definitions Used in Histopathologic Evaluations and an Explanation of the Grading of Lesion Severity

Microblister: Loss of epidermal basal cell attachment to the underlying basement membrane of at least two adjacent cells. The loss of attachment creates a space which may appear empty, full of proteinaceous fluid, or filled with neutrophils. One or a few isolated small areas of detachment is graded 1, minimal. Many such areas of detachment, or several larger (10 or more contiguous cells) areas of detachment is graded 2, mild. When half or more of the epidermis in the tissue section is detached from the dermis, it is graded 3, moderate. Such lesions typically have a much larger space between the basal cells and the dermis. When nearly all of the epidermis is separated from the dermis, it is graded 4, marked. In such situations, there are usually focal, point attachments, so the entire epidermis is not lifted along the full width of the section.

Epidermal necrosis: The epidermal cells exhibit cytoplasmic eosinophilia, nuclear loss or pyknosis, and are generally shrunken. If only individual cells are affected, it is graded 1 (these are generally isolated basal cells). If small areas are affected, with normal areas in close proximity, it is graded 2. If the epidermis exhibits cell death in a full-thickness (all layers of epidermis) pattern, and affects half or more of the skin section, it is graded 3. If the epidermis is virtually entirely necrotic, it is graded 4. Severe ulcers assume that the epidermis is necrotic.

Follicular necrosis: If isolated epithelial cells of the hair follicles exhibit eosinophilia or pyknosis, it is graded 1. If clusters of adjacent cells within follicles are dead, it is graded 2. If cells of half or more of a particular hair follicle are dead, it is graded 3. Grade 4 lesions have complete necrosis of the follicular epithelium underlying much of the epidermal lesion area. This indicates that the agent has penetrated deeply.

Dermal necrosis: Loss of collagen fiber integrity, evidenced by pale eosinophilic staining and homogeneous appearance, indicates necrosis of dermal fibers. With only isolated areas, it is graded 1. Multiple areas are graded 2. Necrosis of most of the superficial dermal collagen in the lesion area is graded 3. A grade four lesion requires deep (to the base of the associated adnexa) dermal necrosis.

Hemorrhage: Extravasated erythrocytes is hemorrhage. A few isolated foci is graded 1. Multiple, common foci is graded 2. Large pools of blood is graded 3. A grade four lesion requires a massive area of blood pooling with displacement of large areas of dermal collagen.

Vascular necrosis: Loss of integrity of a medium to large blood vessel is vascular necrosis. Grading depends upon the number of vessels affected and the severity. Partial necrosis of one vessel is graded 1 to 2. Complete necrosis of a vessel is graded 3; multiple such lesions are graded 4.

Pustular epidermitis: Collections of neutrophils in the epidermis proper is graded by extent; one or two small foci is graded 1; three or more small foci is graded 2; one or more large foci is graded 3; a grade four lesion would indicate massive infiltration of the entire epidermis by neutrophils.

Task 95-38, Phase 2a, Day 1

Key for HGPs #301 and 305 dosed 2/19/1996. Exposure duration - 2 hr.

Animal # 301

Site	Treatment
Α	10 μL of 10% HD in CHCl ₃
В	50 μL of 10% HD in CHCl ₃
С	10 μL of 10% HN in CHCl ₃
D	50 μL of 10% HN in CHCl ₃
E	10 μL of 10% L in CHCl ₃
F	50 μL of 10% L in CHCl ₃
G	1 μL of neat HD
H	

Site	Treatment
A	10 μL of 10% HN in CHCl ₃
В	50 μL of 10% HN in CHCl ₃
С	10 μL of 10% L in CHCl ₃
D	50 μL of 10% L in CHCl ₃
Е	10 μL of 10% HD in CHCl ₃
F	50 μL of 10% HD in CHCl ₃
G	1 μL of neat HD
Н	

Dosing Date: 2/19/96

MREF Task 95-38 G1555-38A

Animal # 301	Site	A	В	С	D	E	F	G
Histopathology Markers:								
Microblister		2	2	2	2	3	4	2
Epidermal Necrosis		2	4	4	3	3	4	3
Follicular Necrosis		3	4	4	4	2	4	4
Dermal Necrosis		0	0	0	0	0	0	0
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	1	2	0
Pustular Epidermitis		0	0	0	0	0	0	0
Notes: all lesions are centrally located; some normal skin preserall	nt on		mild dermal inflam	min dermal inflam	min dermal inflam	mild dermal inflam	mild dermal inflam	min dermal inflam

Animal # 305	Site	A	В	С	D	Е	F	G
Histopathology Markers:								
Microblister		2	2	3	3	3	2	2
Epidermal Necrosis		4	4	4	4	4	4	4
Follicular Necrosis		3	4	4	4	4	4	4
Dermal Necrosis		0	0	0	0 .	0	0	0
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		0	0	1	0	1	1	0
Pustular Epidermitis		0	0	0	0	0	0	0
Notes: all lesions are centrally located; some normal skin presentall	nt on	mild dermal inflam	mild dermal inflam	mild dermal inflam		mild dermal inflam	min dermal inflam	mild dermal inflam

Degree of Severity Grading Scale:

0 = Normal, 1 = Minimal, 2 = Intermediate, 3 = Moderate, 4 = Severe

A. W. Singer, DVM, DACVP

Task 95-38, Phase 2a, Day 2

Key for HGPs #306 and 309 dosed 2/21/1996. Exposure duration - 2 hr.

Animal # 306

Site	Treatment
A	5 μL of 10% L in CHCl ₃
В	10 μL of 10% L in CHCl ₃
С	5 μL of 10% HD in CHCl ₃
D	10 μL of 10% HD in CHCl ₃
E	5 μL of 10% HN in CHCl ₃
F	10 μL of 10% HN in CHCl ₃
G	1 μL of neat HD
Н	

Site	Treatment
Α	10 μL of 10% HD in CHCl ₃
В	5 μL of 10% HD in CHCl ₃
С	10 μL of 10% HN in CHCl ₃
D	5 μL of 10% HN in CHCl ₃
E	10 μL of 10% L in CHCl ₃
F	5 μL of 10% L in CHCl ₃
G	1 μL of neat HD
Н	

Animal # 306	Site	A	В	С	מ	E	F	G
Histopathology Markers:								-
Microblister		4	3	1***	3	4	2***	1***
Epidermal Necrosis		4	4*	4***	4	4	4***	4***
Follicular Necrosis		4	4	4	4	4	4	4
Dermal Necrosis		1	1**	2	0	0	2	2
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		1	0	0	0	0	0	0
Pustular Epidermitis		0	0	0	0	0	0	0
Notes: *focal ulceration **deep dermal edema ***large ulcer precludes much blister potential	5	mod dermal inflam	mild dermal inflam	mild dermal inflam	min dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam

Animal # 309	Site	A	В	С	D	Е	F	G	
Histopathology Markers:									
Microblister		3	0*	4	4	4	4	3	
Epidermal Necrosis		3	4*	4	4	4	4	4	
Follicular Necrosis		4	4	3	2	4	3	4	
Dermal Necrosis		1	2	0	0	0**	0	0	
Vascular Necrosis		0	0	0	0	0	0	0	
Hemorrhage		0	0	0	0	1	0	0	
Pustular Epidermitis		1	0	1	1	0	0	0	
Notes: *large ulceration preclud blister potential **deep dermal edema	les	mild dermal inflam	mild dermal inflam	mod dermal inflam	mod derm infla m	mod dermal inflam	mod dermal inflam	mild derm infla m	

Note: Some normal skin is present on all sections, both animals; lesions are centrally located in trimmed area.

Degree of Severity Grading Scale:

0 = Normal; 1 = Minimal; 2 = Mild; 3 = Moderate; 4 = Severe

Allen W. Singer, D.V.M.

Task 95-38, Phase 2a, Day 3

Key for HGPs #312 and 316 dosed 2/27/1996. Exposure duration - 1 hr.

`Animal # 312

Site	Treatment
A	10 μL of 10% L in CHCl ₃
В	5 μL of 10% L in CHCl ₃
С	10 μL of 10% HD in CHCl ₃
D	5 μL of 10% HD in CHCl ₃
E	10 μL of 10% HN in CHCl ₃
F	5 μL of 10% HN in CHCl ₃
G	1 μL of neat HD
H	ì

Site:	Treatment
A	10 μL of 10% HN in CHCl ₃
В	5 μL of 10% HN in CHCl ₃
С	10 μL of 10% L in CHCl ₃
D	5 μL of 10% L in CHCl ₃
Е	10 μL of 10% HD in CHCl ₃
F	5 μL of 10% HD in CHCl ₃
G	1 μL of neat HD
Н	

Animal # 312	Site	A	В	С	D	E	F	G
Histopathology Marker	s:							
Microblister	3	3	3	3	4	3	3	
Epidermal Necrosis		4	4	4	4	4	4	4
Follicular Necrosis		4	4	4	4	4	3	4
Dermal Necrosis		0*	0*	0	0**	0	0	0*
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		1	2	0	0	0	0	0
Pustular Epidermitis		0	0	0	0	1	2	0
Notes: *mod dermal edem **minimal dermal edem		mild dermal inflam	mod dermal inflam	mild dermal inflam	mild derm infla m	mild dermal inflam	mod derm inflam	mild derm infla m
Animal # 316	Site	A	В	С	D	Е	F	G
Histopathology Markers	s:			;			,	·
Microblister		3	4	4	4	3	3	3
Epidermal Necrosis		44	4	4 .	4	4	4	4
Follicular Necrosis		4	3	4	4	4	4	4
Dermal Necrosis		0*	0	0**	0**	0	1	0**
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		0	0	2	2	1	0	0
Pustular Epidermitis		1	1	0	0	1	1	2
Notes: *minimal dermal edema **moderate dermal eden	na	mod dermal inflam	mod dermal inflam	mod dermal inflam	sever e derm infla m	mild dermal inflam	mod dermal inflam	mod derm infla m

Note: All sections (312 and 316) have normal, unaffected skin at one or both margins of the section.

Degree of Severity Grading Scale:

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

Allen W. Singer, D.V.M.
Appendix E

Task 95-38, Phase 2b, Day 1

Key for HGPs #311, 313, 315, 317, and 324 dosed 3/5/1996. Exposure duration - 1 hr.

Animal #311

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	20 μL of neutralization solution
С	5 μL of 10% L in CHCl ₃
D	20 μL of neutralization solution
E	5 μL of 10% HD in CHCl ₃
F	20 μL of neutralization solution
G	1 μL of neat HD
H	20 μL of neutralization solution

Site	Treatment
A	5 μL of 10% HD in CHCl ₃
В	20 μL of neutralization solution
С	5 μL of 10% HN in CHCl ₃
D	20 μL of neutralization solution
Е	5 μL of 10% L in CHCl ₃
F	20 μL of neutralization solution
G	1 μL of neat HD
H	20 μL of neutralization solution

Animal # 315

Site	Treatment
Α	5 μL of 10% HN in CHCl ₃
В	20 μL of neutralization solution
С	5 μL of 10% L in CHCl ₃
D	20 μL of neutralization solution
E	5 μL of 10% HD in CHCl ₃
F	20 μL of neutralization solution
G	1 μL of neat HD
Н	20 μL of neutralization solution

Site	Treatment
A	5 μL of 10% L in CHCl ₃
В	20 μL of neutralization solution
С	5 μL of 10% HD in CHCl ₃
D	20 μL of neutralization solution
E	5 μL of 10% HN in CHCl ₃
F	20 μL of neutralization solution
G	1 μL of neat HD
Н	20 μL of neutralization solution

Site	Treatment
A	5 μL of 10% HD in CHCl ₃
В	20 μL of neutralization solution
С	5 μL of 10% HN in CHCl ₃
D	20 μL of neutralization solution
E	5 μL of 10% L in CHCl ₃
F	20 μL of neutralization solution
G	1 μL of neat HD
H	20 μL of neutralization solution

E-11

Animal # 311	Site	A	В	С	D	E	F	G	Н
Histopathology Markers:									
Microblister		3	0	3	0	2	0	3	0
Epidermal Necrosis		4	0	4	0	4	0	4	0
Follicular Necrosis		2	0	4	0	4	0	4	0
Dermal Necrosis		0	0	0	0	0	0	0*	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		2	0	3	0	0	0	1	0
Pustular Epidermitis		2	0	0	0	11	0	0	0
Note: *moderate deep dermal edema		mod dermal inflam		mod dermal inflam		mod dermal inflam		mod derm infla m	

Animal # 313	Site	A	В	С	D	E	F	G	Н
Histopathology Markers:									
Microblister		3	0	4	0	4	0_	2	0
Epidermal Necrosis		4	0	4	0	4	0	4	0
Follicular Necrosis		4	0	4	0	3	0	4	0
Dermal Necrosis		0	0	1	0	0	0	0*	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	1	0	0	0
Pustular Epidermitis		0	0	0	0	0	0	0	٥
Note: *moderate deep dermal edema		mild derm inflam		mod dermal inflam	min dermal inflam	mod dermal inflam		mild dermal inflam	

Animal # 315	Site	A	В	С	D	Е	F	G	H
Histopathology Markers:									
Microblister		2	0	4	0	3	0	2	0
Epidermal Necrosis		3	0	4	0	4	0	4	0
Follicular Necrosis		2	0	4	0	4	0	4	0
Dermal Necrosis		0	0	1	0	1	0	0*	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		0	0	1	0	2	0	0	0
Pustular Epidermitis		1	0	0	0	0	0	0	0
Note: *moderal dermal edema		mod dermal inflam		marked dermal inflam		mod dermal inflam		mild dermal inflam	

Animal # 317	Site	A	В	С	D	Е	F	G	н
Histopathology Markers:									
Microblister		2	0	2	0	3	0	2	0
Epidermal Necrosis		4	0	4	0	4	0	4	0
Follicular Necrosis		4	0	4	0	3	0	4	0
Dermal Necrosis		0*	0	2**	0	0	0	0*	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Нетогтнаде		2	0	1	0	0	0	0	0
Pustular Epidermitis		0	0	0	0	2	0 0		0
Notes: *mild dermal edema **focal ulceration(s)		mild dermal inflam	min dermal inflam	mod dermal inflam		mod dermal inflam		mild dermal inflam	

E-13

	7	7	7	7	T		7	7	7
Animal # 324	Site	A	В	C	D	E	F	G	H
Histopathology Markers:									
Microblister		4	0	4	0	4	0	3	0
Epidermal Necrosis		4	0	4	0	4	0	4	0
Follicular Necrosis		4	0	2	0	4	0	4	0
Dermal Necrosis		1	0	0	0	0	0	0	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	1	0	0	0
Pustular Epidermitis		0	0	1	0	0	0	0	0
Notes:		mod dermal inflam	min dermal inflam	mod dermal inflam		mod dermal inflam		min dermal inflam	

Note: Normal (unaffected) skin present laterally on all sections where lesions were observed.

Histopathological Markers Degree of Severity Grading Scale DVM

3/7/96 Allen W. Singer,

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

Task 95-38, Phase 3, Day 1

Key for HGPs #310, 491, 493, and 498 dosed 3/13/1996. Exposure duration - 1 hr.

Animal # 310

Site	Treatment						
A	5 μL of 10% HD in CHCl ₃						
В	25 μL of Red waste stream						
С	5 μL of 10% HN in CHCl ₃						
D	25 μL of Blue waste stream						
Е	5 μL of 10% L in CHCl ₃						
F	25 μL of Charcoal waste stream						
G	1 μL of neat HD						
H	i						

Site	Treatment						
Α	5 μL of 10% L in CHCl ₃						
В	25 μL of Charcoal waste stream						
С	5 μL of 10% HD in CHCl ₃						
D	25 μL of Red waste stream						
Е	5 μL of 10% HN in CHCl ₃						
F	25 μL of Blue waste stream						
G	1 μL of neat HD						
H							

E-15

Site	Treatment						
Α	5 μL of 10% HD in CHCl ₃						
В	25 μL of Red waste stream						
С	5 μL of 10% HN in CHCl ₃						
D	25 μL of Blue waste stream						
Е	5 μL of 10% L in CHCl ₃						
F	25 μL of Charcoal waste stream						
G	1 μL of neat HD						
Н							

Site	Treatment					
A	5 μL of 10% HN in CHCl ₃					
В	25 μL of Blue waste stream					
С	5 μL of 10% L in CHCl ₃					
D	25 μL of Charcoal waste stream					
E	5 μL of 10% HD in CHCl ₃					
F	25 μL of Red waste stream					
G	1 μL of neat HD					
Н						

E-16

								
Animal # 310	Site	A	В	C	D	E	F	G
Histopathology Marl	cers:							
Microblister		2	0	4	2	1	0	1
Epidermal Necrosis		4	1	4	4	4*	2	4*
Follicular Necrosis		4	0	4	1	4	0	4
Dermal Necrosis		0	0	1	0	3	0	3**
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	0	0	1
Pustular Epidermitis		0	1	0	1	0	1	0
Notes: *marked ulceration **moderate dermal edema		mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam	mild dermal inflam	mod dermal inflam
		<u> </u>		; 			1	
Animal # 491	Site	A	В	С	D	E	F	G
Histopathology Mark	ers:		_r					,
Microblister		4	0	1	0	4	3	2
Epidermal Necrosis		4*	1	4**	0	4	4	4
Follicular Necrosis	· 	4	0	4	0	3	0	4
Dermal Necrosis		3	0	3	0	0	0	0
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		2	0	0	0	0	0	0
Pustular Epidermitis		0	0	0	0	0	0	0
Notes: *mild ulceration **marked ulceration	n	mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam

					_									
Animal # 493	Animal # 493 Site			В		С		D		E		F		G
Histopathology Mark	cers:													
Microblister	Microblister			0		4		4		2		0		2*
Epidermal Necrosis		4**		0		4		4		4**		1		4**
Follicular Necrosis		4		0		3		0		4		0		4
Dermal Necrosis		3		0		0		0		3		0		3
Vascular Necrosis		0		0		0		0		0		0		0
Hemorrhage		0		0		0		0		1	\neg	0		0
Pustular Epidermitis		0		1		0		0		0		0		0
Notes: *at edge of ulcer **marked ulceration		mod derma inflan	al	mod dermal inflam		mod derma inflan	ıl	mod dermal inflam		mod dermal inflam		mod dermal inflam		mod dermal inflam
Animal # 498	Site	A		В		C :		D		E		F		G
Histopathology Mark	ers:		,											
Microblister		2*		3	Ŀ	3		0		3	L	0		1
Epidermal Necrosis		4**	4	ţ***	Ĺ	4***		0	4	**		0		4***
Follicular Necrosis		4		0		4		0		4		0		4
Dermal Necrosis		3		1		2		0		3		0		2
Vascular Necrosis		0		0		0		0		0		0		0
Hemorrhage		0		0		1		0		0		0		0
Pustular Epidermitis		1		0		0		1		1		0		0
Notes: *at edge of ulcer **marked ulceration ***minimal ulceration		mod dermal inflam	de	nild ermal flam	đ	mod ermal ıflam	de	nod rmal flam	der	od mal lam	de	nild rmal flam	d	mild ermal nflam

Histopathological Markers: Degree of Severity Grading Scale 3/18/96 0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe Allen W. Singer, DVM

Task 95-38, Phase 3, Day 2

Key for HGPs #494, 496, 497, and 499 dosed 3/21/1996. Exposure duration - 1 hr.

Animal # 494

Site	Treatment						
A	5 μL of 10% L in CHCl ₃						
В	25 μL of Charcoal waste stream						
С	5 μL of 10% HD in CHCl ₃						
D	25 μL of Red waste stream						
Е	5 μL of 10% HN in CHCl ₃						
F	25 μL of Blue waste stream						
G	1 μL of neat HD						
H							

Site	Treatment
Α	5 μL of 10% HD in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Blue waste stream
Е	5 μL of 10% L in CHCl ₃
F	25 μL of Charcoal waste stream
G	1 μL of neat HD
Н	

E-19

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Charcoal waste stream
E	5 μL of 10% HD in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
Н	

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Charcoal waste stream
E	5 μL of 10% HD in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
н	

E-20

Animal # 494	Site	A	В	С	D	E	F	G			
Histopathology Markers:											
Microblister		4	0	1	0	3	2	3			
Epidermal Necrosis		4	0	4**	0	4	2	4***			
Follicular Necrosis		4	0	4	0	4	0	4			
Dermal Necrosis		0*	0	3	0	0	0	2			
Vascular Necrosis		0	0	0	0	0	0	0			
Hemorrhage		3	0	0	0	0	Ō	0			
Pustular Epidermitis		0	0	1	0	1	0	0			

Animal # 496	Site	A	В	С	D	E	F	G		
Histopathology Markers:										
Microblister		0	0	0	4	1	0	2		
Epidermal Necrosis		4*	0	4*	3	4*	1	4*		
Follicular Necrosis		4	0	4	0	4	0	4		
Dermal Necrosis		3	0	3	0	4	0	3**		
Vascular Necrosis		0	0	0	0	0	0	0		
Hemorrhage		0	0	0	0	0	0	0		
Pustular Epidermitis		0	0	0	0	0	0	0		
Notes: *marked ulco precludes potential blister **mild dermal edem		mod dermal inflam	min dermal inflam	mod dermal inflam	mild dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam		

Animal # 497 Site		A	В		С	D	E		F	G
Histopathology Mari	kers:									
Microblister	1	2		4	0	2		0	2	
Epidermal Necrosis		4*	4		4	1***	* 4*		0	4*
Follicular Necrosis		4	0		4	0	4		0	4
Dermal Necrosis		3	0		0**	0	2		0	2**
Vascular Necrosis		0	0		0	0	0		0	0
Hemorrhage		0	0	!	0	0	0		0	0
Pustular Epidermitis		0	0		0	0	0		0	0
Notes: *marked ulcer **moderate dermal e ***mild epithelial cel	dema	mod derma inflan	dern	ıal	mod derma inflan	derma	1	al	mild dermal inflam	
Animal # 499	Site	A	В		С	D	Е		F	G
Histopathology Mark	ers:									
Microblister		4	2		3	0	4		0	3
Epidermal Necrosis		4	3		4	2	4		0	4
Follicular Necrosis		4	0		4	0	4		0	4
Dermal Necrosis	Dermal Necrosis		0		2*	0	2		0	1*
Vascular Necrosis		0	0		0	0	0		0	0
Hemorrhage	Hemorrhage		0		1	ο	0		0	0
Pustular Epidermitis		1	0		0	0	1		0	0
Note: *mild dermal ed		mod dermal inflam	mild dermal inflam	d	mod ermal nflam	min dermal inflam	mod dermal inflam	- 1	min Iermal inflam	mild dermal inflam

Histopathological Markers: Degree of Severity Grading Scale 3/25/96 0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe Allen W. Singer, DVM

Task 95-38, Phase 3, Day 3

"Fresh" Blue and Red waste streams received 6/19/1996

Key for HGPs #339, 341, 342, and 346 dosed 6/20/1996. Exposure duration - 1 hr.

Animal # 339

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	25 μ L of Red waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Blue waste stream
Е	5 μL of 10% HD in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
Н	$25 \mu L$ of Blue waste stream

Site	Treatment
Α	5 μL of 10% HD in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Red waste stream
E	5 μL of 10% L in CHCl ₃
F	25 μL of Blue waste stream
G	1 μL of neat HD
Н	25 μL of Red waste stream

E-23

Site	Treatment
Α	5 μL of 10% L in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Red waste stream
E	5 μL of 10% HN in CHCl ₃
F	25 μL of Blue waste stream
G	1 μL of neat HD
Н	25 μL of Red waste stream

Site	Treatment
Α	5 μL of 10% L in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Blue waste stream
E	5 μL of 10% HN in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
Н	25 μL of Blue waste stream

Animal # 339	Site	A	В	С	D	Е	F	G	H		
Histopathology Mark	Histopathology Markers:										
Microblister		3	0	4	3	3	0	2	2		
Epidermal Necrosis		4	0	4	4	4**	0	4	2		
Follicular Necrosis		4	0	4	0	4	0	4	0		
Dermal Necrosis		0	0	2*	0	2	0	0*	0		
Vascular Necrosis		0	0	0	0	0	0	0	0		
Hemorrhage		0	0	0	0	0	0	0	0		
Pustular Epidermitis		1	0	1	1	1	0	0	0		
Notes: *moderate der edema **focal ulceration(s)	mal	mod dermal inflam	min dermal inflam	mod dermal inflam	mild dermal inflam	mod dermal inflam	min dermal inflam	mild dermal inflam	min dermal inflam		

Animal # 341	Site	A	В	С	D	Е	F	G	Н	
Histopathology Markers:										
Microblister		2	2	2	. 0	3	0	2	0	
Epidermal Necrosis	1818	4*	4	4*	0	4*	4*	4*	0	
Follicular Necrosis		4	1	4	0	4	2	2	0	
Dermal Necrosis		3	1	2	0	3**	3	3**	0	
Vascular Necrosis		0	0	0	0	0	0	0	0	
Hemorrhage		1	0	0	0	1	0	1	0	
Pustular Epidermitis		0	0	0	0	0	0	0	0	
Notes: *focal ulceration(**moderate dermal eden		mild dermal inflam	mild dermal inflam	mild dermal inflam	min dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam	min dermal inflam	

Animal # 342	Site	A	В	С	D	E	F	G	H
Histopathology Markers	:								
Microblister		3	1	3	0	4	3	3	1
Epidermal Necrosis		4	4	4	0	4	4	4	4
Follicular Necrosis		4	0	4	0	4	1	4	4
Dermal Necrosis		0*	0	0*	0	0	0	0	0*
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	0	0	0	0
Pustular Epidermitis		0	0	1	0	0	0	0	0
Notes: *mild to moderate dermal edema	•	mild dermal inflam	min dermal inflam	mild dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	mild dermal inflam	min dermal inflam

Animal # 346	Site	Α	В	С	D	Е	F	G	H	
Histopathology Markers:										
Microblister		2	0	2	1	4	0	2	1	
Epidermal Necrosis		4	0	4	4	4	0	4	4**	
Follicular Necrosis		4	0	4	1	4	0	4	0	
Dermal Necrosis		0*	0	0	0	2	0	0*	0	
Vascular Necrosis		0	0	0	0	0	0	0.	0	
Hemorrhage		0	0	0	0	0	0	0	0	
Pustular Epidermitis		0	0	0	0	0	0	0	0	
Notes: *moderate derma edema; **most of surface epitheli artifactually stripped awa	ium	mild dermal inflam	min dermal inflam	mod dermal inflam	mild dermal inflam	mild dermal inflam		mild dermal inflam	min dermal inflam	

Note: Normal (unaffected) skin presented laterally on all skin sections with lesions.

Histopathological Markers

6/25/96

Degree of Severity Grading Scale

Allen W. Singer, DVM

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

Task 95-38, Phase 3, Day 4

[&]quot;Fresh" Blue and Red waste streams received 6/19/1996
Appendix E 229

E-26

Key for HGPs #340, 345, 351, and 352 dosed 6/26/1996. Exposure duration - 1 hr.

Animal # 340

Site	Treatment
Α	5 μL of 10% HD in CHCl ₃
В	25 μ L of Blue waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Red waste stream
Е	5 μL of 10% L in CHCl ₃
F	$25 \mu L$ of Blue waste stream
G	1 μL of neat HD
Н	25 μL of Red waste stream

Site	Treatement
A	5 μL of 10% L in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Blue waste stream
Е	5 μL of 10% HN in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
H	25 μL of Blue waste stream

Site	Treatment
Α	5 μL of 10% HN in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Red waste stream
E	5 μL of 10% HD in CHCl ₃
F	25 μL of Blue waste stream
G	1 μL of neat HD
Н	25 μL of Red waste stream

Site	Treatment
Α	5 μL of 10% HD in CHCl ₃
В	25 μ L of Red waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Blue waste stream
E	5 μL of 10% L in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
H	$25~\mu L$ of Blue waste stream

Animal # 340	Site	Α	В	С	D	E	F	G	Н	
Histopathology Ma	Histopathology Markers:									
Microblister		2*	3	3	0	3	3	0	0	
Epidermal Necro	sis	4**	4	4	0	4	4	4	1	
Follicular Necrosis		4	2	4	0	4	0	4	0	
Dermalal Necrosis		2	0	1	0	0***	0	3***	0	
Vascular Necrosis		0	0	0	0	0	0	0	0	
Hemorrhage		2	0	0	. 0	2	0	2	0	
Pustular Epidermi	is	0	0	0	0	0	0	0	0	
Notes: *at edge of t **mild ulceration ***mild dermal . e		mod dermal inflam	mild dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	min dermal inflam	min dermal inflam	

Animal # 345	Site	Α	В	С	D	E	F	G	H	
Histopathology Mar	Histopathology Markers:									
Microblister		3*	0	_2	1	1	0	1	2	
Epidermal Necrosis		4**	0	4	4	4**	0	4	4	
Follicular Necrosis		3	0	4	1	4	0	4	1	
Dermal Necrosis		3	0	0	0	3	0	2***	0	
Vascular Necrosis		0	0	0	0	0	0	0	0	
Hemorrhage		2	0	1	0	2	0	1	0	
Pustular Epidermit	is	0	1	0	0	0	0	0	0	
Notes: *at one edge ulcer **marked ulceration present ***mild dermal ed		mod dermal inflam	min dermal inflam	mild dermal inflam	mild dermal inflam	mod dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam	

Animal # 351	Site	A	В	С	D	Е	F	G	Н	
Histopathology Marl	Histopathology Markers:									
Microblister		1	2	4	0	1*	2	1	0	
Epidermal Necrosi	s	4	4	4	0	4**	4	4	0	
Follicular Necrosis		4	1	3	0	4	1	4	0	
Derma! Necrosis		0	0	0	0	3	0	3	0	
Vascular Necrosis		0	0	0	0	0	0	0	0	
Hemorrhage		0	0	1	0	1	0	1	0	
Pustular Epidermit:	İs	0	0	0	0	0	0	0	0	
Notes: *at one edge ulcer **marked ulceration present	of	mild dermal inflam	min dermal inflam	mod dermal inflam		mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	

Animal # 352	Site	A	В	С	D	E	F	G	н
Histopathology Markers:									
Microblister		1*	0	2	1	3	0	2	2
Epidermal Necros	sis	4**	0	4**	4	4	0	4**	4
Follicular Necrosis		4	0	3	0	4	0	4	1
Dermal Necrosis		2	0	1	0	0***	0	3	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		2	0	1	0	1	0	1	0
Pustular Epidermit	is	0	0	1	1	0	0	0	0
Notes: *at edge of u **moderate ulcerati ***mild dermal ed	on	mod dermal inflam	min dermal inflam	mod dermal inflam	mod dermal inflam	mild derma l inflam		mod dermal inflam	mild dermal inflam

Histopathological Markers: Degree of Severity Grading Scale 7/1/96

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe Allen W. Singer, DVM

Task 95-38, Phase 3, Day 5

Blue and Red waste streams received 11/28/1995; Charcoal waste stream received 1/25/96.

Appendix E 233

E-30

Equal volumes of waste streams and 10% HD, HN and L solutions - 10 μ L

Key for HGPs #383, 385, 389, and 400 dosed 8/13/1996. Exposure duration - 1 hr.

Animal # 383

Site	Treatment
A	10 μL of 10% L in CHCl ₃
В	$10 \mu L$ of Blue waste stream
С	10 μL of 10% HD in CHCl ₃
D	10 μL of Charcoal waste stream
E	10 μL of 10% HN in CHCl ₃
F	$10 \mu L$ of Red waste stream

Site	Treatment
Α	10 μL of 10% HN in CHCl ₃
В	10 μL of Red waste stream
С	10 μL of 10% L in CHCl ₃
D	10 μL of Blue waste stream
E	10 μL of 10% HD in CHCl ₃
F	10 μL of Charcoal waste stream

Site	Treatment
A	10 μL of 10% HN in CHCl ₃
В	10 μL of Red waste stream
С	10 μL of 10% L in CHCl ₃
D	10 μL of Blue waste stream
Е	10 μL of 10% HD in CHCl ₃
F	10 μL of Charcoal waste stream

Site	Treatment
Α	10 μL of 10% HD in CHCl ₃
В	10 μL of Charcoal waste stream
С	10 μL of 10% HN in CHCl ₃
D	10 μL of Red waste stream
E	10 μL of 10% L in CHCl ₃
F	10 μL of Blue waste stream

E-32 MREF Task 95-38 G1555-38A

Animal # 383 Site		A	В	С	D	Е	F
Histopathology Markers:							
Microblister		3	0	2	0	3	0
Epidermal Necrosis		4	0	4	0	4	0
Follicular Necrosis		4	0	4	0	3	0
Dermal Necrosis		0*	0	0	0	0	0
Vascular Necrosis		0	0	0	0	0	0
Hemorrhage		2	0	1	0	0	0
Pustular Epidermitis		0	1	0	0	1	0
Notes: *moderate dermal edema		mod dermal inflam		mild dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam
Animal # 385	Site	A	В	С	D	Е	F
Histopathology Markers:							
Microblister		4	0	4	1	3	0
Epidermal Necrosis		4	0	4	1**	4	0
Follicular Necrosis		4	0	4	0	4	0
Dermal Necrosis		1	0	0*	0	0*	0
Vascular Necrosis		0	0	0	0	0	0
Нетоптаде		0	0	2	0	0	0
Pustular Epidermitis		l	0	0	0	0	0
Notes: *mod dermal edema **vacuolar degeneration of epith cells leading to intra- and subepithelial microblister		marked dermal inflam	min dermal inflam	mod dermal inflam		mild dermal inflam	min dermal inflam

Histopathological Markers: Degree of Severity Grading Scale 0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

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E-33 MREF Task 95-38 61555-38a

В

Site

Α

Animal # 389

Histopathology Markers:							
Microblister	3	0	2	2	2	0	
Epidermal Necrosis		4	0	4	2	4	1
Follicular Necrosis		2	0	4	1	4	0
Dermal Necrosis		0	0	0*	0	0*	0
Vascular Necrosis		0	0	0	0	0	0
Hemorrhage		1	0	3	0	2	0
Pustular Epidermitis		1	0	0	0	0	1
Notes: *severe dermal edema		mod dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam
Animal # 400 S	ite	A	В	С	D	E	F
Histopathology Markers:				•	<u> </u>		
Microblister		3	0	4	0	3	3
Epidermal Necrosis		4	0	4	0	4	2
Follicular Necrosis		4	0	2	0	4	1
Dermal Necrosis		0*	0	0*	0	0**	. 0
Vascular Necrosis		0	0	0	0	1	0
Hemorrhage	0	0	1	0	3	1	
Pustular Epidermitis	1	0	1	.0	0	0	
Notes: *mild dermal edema **severe dermal edema	mod dermal inflam		mod dermal inflam	min dermal inflam	mod dermal inflam	mild dermal inflam	

Note: Some normal (unaffected) skin present at one or both ends of each section where lesions were present.

Histopathological Markers: Degree of Severity Grading Scale
0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

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Task 95-38, Phase 3, Day 6

"Fresh" Charcoal waste stream received 8/29/96.

 $25~\mu L$ of freshly prepared Charcoal waste stream and 5 μL of 10% HD, HN and L solutions

Key for HGPs #379, 380, 387, and 388 dosed 8/29/1996. Exposure duration - 1 hr.

Animal # 379

Site	Treatment
A	5 μL of 10% L in CHCl ₃
В	25 μL of Charcoal waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Charcoal waste stream
Е	5 μL of 10% HN in CHCl ₃
F	25 μL of Charcoal waste stream

Site	Treatment			
A	5 μL of 10% HN in CHCl ₃			
В	25 μL of Charcoal waste stream			
С	5 μL of 10% L in CHCl ₃			
D	25 μL of Charcoal waste stream			
Е	5 μL of 10% HD in CHCl ₃			
F	25 μL of Charcoal waste stream			

Site	Treatment				
A	5 μL of 10% HD in CHCl ₃				
В	25 μL of Charcoal waste stream				
С	5 μL of 10% HN in CHCl ₃				
D	25 μL of Charcoal waste stream				
Е	5 μL of 10% L in CHCl ₃				
F	25 μL of Charcoal waste stream				

Site	Treatment
Α	5 μL of 10% L in CHCl ₃
В	25 μL of Charcoal waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Charcoal waste stream
Е	5 μL of 10% HN in CHCl ₃
F	25 μL of Charcoal waste stream

E-36

MREF Task 95-38 G1555-38A

Animal # 379	Site	Α	В	С	D	Е	F
Histopathology Markers							
Microblister		4	0	2	0	3	0
Epidermal Necrosis		4	1**	4	1**	4	1**
Follicular Necrosis		4	1**	4	1**	4	I**
Dermal Necrosis		0	0	0*	0	0	0
Vascular Necrosis		0	0	0	0	0	0
Нетоптаде		2	0	2	0	1	0
Pustular Epidermitis		0	0	0	0	0	0
Notes: *moderate dermal edema; **random single-cell necrosis noted		mod dermal inflam	min dermal inflam	mild dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam
Animal # 380	Site	A	В	С	D	Е	F
Histopathology Markers:							
Microblister		4	0	4	0	3	0
Epidermal Necrosis		4	l*	4	1*	4	0
Follicular Necrosis		4	1*	4	1*	4	1*
Dermal Necrosis		1	0	2**	0	3**	0
Vascular Necrosis		0	0	0	0	0	0
Hemorrhage	0	0	1	0	0	0	
Pustular Epidermitis	0	0	0	0	0	0	
Notes: *random single cell necrosis **mod dermal edema; focal ulcer in area of necrosis		mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam

Note: Some normal (unaffected) skin present at one or both ends of each section where lesions were present. Histopathological Markers: Degree of Severity Grading Scale 9/9/96

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

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E-37 MREF Task 95-38 1555-38-A

Animal # 387 Site		Α	В	С	D	E	F
Histopathology Markers:							
Microblister		3	0	2	0	4	0
Epidermal Necrosis		4	1**	4	1**	4	1**
Follicular Necrosis		4	1**	3	1**	4	1**
Dermal Necrosis		0	0	0	0	0*	0
Vascular Necrosis		0	0	0	0	0	0
Hemorrhage		0	0	0	0	2	0
Pustular Epidermitis		0	0	0	0	0	1
Notes: *moderate dermal edema; **random single-cell necrosis noted		mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam
Animal # 388 Site		Α	В	С	D	Е	F
Histopathology Markers	:						
Microblister		4	0	4	0	3	0
Epidermal Necrosis		4	1*	4	1*	4	1*
Follicular Necrosis		4	1*	4	1*	2	1*
Dermal Necrosis		0**	0	0**	0	0	0
Vascular Necrosis		0	0	0	0	0	0
Hemorrhage		3	0	1	0	0	0
Pustular Epidermitis		0	0	0	0	1	0
Notes: *random suigle cell necrosis **mod dermal edema		mild dermal inflam	min dermal inflam	mild dermal inflam	mild dermal inflam	mod dermal inflam	min dermal inflam

Histopathological Markers: Degree of Severity Grading Scale 9/9/96

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe Allen W. Singer, DVM

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